# ANHARMONIC OSCILLATORS IN QUANTUM MECHANIC

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## ANHARMONIC OSCILLATORS IN QUANTUM MECHANIC

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#### CERTIFICATE

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### CONTENTS

List of	rable	ES	V
List of	Figur	res	vii
bynopsis	3		viii
Chapter	I	Introduction	1
Chapter	II	The Quartic Anharmonic Oscillator	13
		l. Scaling and the Appropriate Scaling Formula	13
		2. Hethod	16
		5. Eigenvalues	20
-		4. Stability of Zeros of $\Delta_m(E)$	22
		5. Checks for the Eigenvalues	26
		6. Eigenfunctions	27
		7. leatures of the Method	30
Chapter	III	The General Anharmonic Oscillator	48
		1. Introduction	48
		2. The Appropriate Scaling Formula	51
		3. Method	55
		4. Eigenvalues	56
Chapter	IV	The Double Minimum Oscillator	68
		1. Introduction	68
		2. Eigenvalues	70
		J. The WKB Formula for Splitting	72
Chapter	V	The Irensition homents	81
		1. The Recurrence Relation in $\langle n   x^k   n' \rangle$	81
		2. Computation of the gransition homents	84
Appendia	c A		\$2
Appendiz	c B		93
Referenc	ces		95

## LIST OF TABLES

Table		Page
I.l	Comparison of the results of various variational	
	and numerical methods.	7
II.l	Stabilization of the computed eigenvalues.	35
11.2	Comparison of our results for $\mathbb{F}_n$ ( $\lambda=1$ ) with results	
	of earlier calculations.	37
II.3	Eigenvalues of the pure quartic oscillator and the	
	quartic anharmonic oscillator for $\lambda = 1$ .	38
II.4	The quartic anharmonic oscillator eigenvalues	
	for various values of $\lambda$ .	41
II.5	Bigenvalues of the quartic anharmonic oscillator	
	in regimes of extreme values of $(n,\lambda)$ .	45
III.l	The sextic anharmonic oscillator eigenvalues for	
	various values of $\lambda$ .	58
I]I.2	The octic anharmonic oscillator engenvalues for	
	various values of $\lambda$ .	63
IV.1	The double minimum oscillator eigenvalues for	
	various values of $\lambda$ .	75
IV.2	Comparison of the WKB values for the splitting with	
	the corresponding accurate values.	80
V.l	The ratio of $[/]$ for the quartic	
	anharmonic and the pure quartic oscillators	87

rable		Page
V.2	The nonzero matrix elements $\langle n x n'\rangle$ and $\langle n x^2 n'\rangle$	
	between the lowest ten states of the pure quartic	
	oscillator for $\lambda = 1$ .	88
v.3	The nonzero matrix elements $\langle n x n'\rangle$ and $\langle n x^2 n'\rangle$	
	between the lowest ten states of the quartic	
	anharmonic oscillator for $\lambda = 1$ .	90

## LIST OF FIGURES

Figure		Page
II.l	The quartic anharmonic oscillator eigenfunc-	
	tions.	47
IV.l	The double minimum oscillator eigenfunctions.	79

#### SYNOPSIS

Thesis entitled: ANHARMONIC OSCILLATORS IN QUANTUM MECHANICS, submitted by SUBODY P. BHATNAGAD to the Department of Physics, Indian Institute of Technology Kanpur, in partial fulfilment of the requirement of the Ph.D. degree.

In this work the following eigenvalue problems have been investigated:

- (i) the quartic anharmonic oscillator (Hamiltonian  $H=p^2+x^2+\lambda x^4$ ,  $p=-i\frac{d}{dx}$ ,  $\lambda$  >0) and the associated problem of the pure quartic oscillator ( $H=p^2+\lambda x^4$ ,  $\lambda$  >0),
- (ii) the general anharmonic oscillators (H =  $p^2 + x^2 + \lambda x^{2\mu}$ ,  $\lambda > 0$ ,  $\mu = 3, 4, ...$ ),
- (iii) the double minimum oscillator (H =  $p^2 x^2 + \lambda x^4$ ,  $\lambda > 0$ ).

Eigenvalues and eigenfunctions of these systems are obtained in all regimes of the quantum number n and the anharmonicity constant  $\lambda$ . The computed eigenfunctions are then used to obtain the transition moments. The eigenvalues reported in this work are accurate to 15 significant figures and the transition moments to 12 figures.

The eigenvalues of the anharmonic oscillators  $(H = p^2 + x^2 + \lambda x^{2\mu})$  fall into two distinct classes (Hioe et al. 1976). In the low n, low  $\lambda$  regime the eigenvalues differ slightly from the harmonic oscillator levels whereas in the high n, high  $\lambda$  regime they differ slightly from the

pure anharmonic oscillator (H =  $p^2 + \lambda x^{2\mu}$ ) eigenvalues. Between these two regimes lies the 'boundary layer' in which the eigenvalues are neither 'near harmonic' nor near 'pure anharmonic'. The existence of different regimes implies different oscillation properties of the corresponding eigenfunctions. We assert that this fact must be explicitly included in solving the eigenvalue problem. The method applied in this work (Banerjee 1976) involves the use of an appropriately (according to regime) scaled basis for the expansion of each eigenfunction. The appropriately scaled basis simulates the different oscillation properties of the eigenfunctions in different regimes and makes possible a uniform treatment of the problem in all regimes.

The Chapter I of the thesis is a review of various earlier methods used to solve the anharmonic oscillator eigenvalue problem. These methods are suitable only in a particular regime of  $(n,\lambda)$  and do not give eigenvalues to the same accuracy when extended to other regimes. The construction of an appropriately scaled basis and the method used in this thesis for the computation of the eigenvalues is described in Chapter II. The actual computation of the eigenvalues is reduced to the determination of the roots of a transcendental equation in the energy. This is done numerically. Accurate eigenvalues and eigenfunctions of the quartic anharmonic and the pure quartic oscillator are

then obtained for various values of  $(n, \lambda)$  covering all different regimes. In Chapter III we show that the eigenvalue problem of the general anharmonic oscillators may be solved for any  $\mu$  in all regimes of  $(n,\lambda)$  using the same method. In Chapter IV accurate eigenvalues and eigenfunctions of the double minimum oscillator are calculated. A WKB expression for the splitting between the lover eigenvalues, bunched in pairs, is obtained and the WKB values are compared with the corresponding accurate values. In Chapter V, the transition moments which are the matrix elements of  $x^k$  (k = integer) between the anharmonic oscillator digenstates are calculated using the computed eigenvalues and eigenfunctions. transition moments for any particular transition satisfy an exact linear recurrence relation (Bancajce 1977) from which the higher moments for that transition may be obtained recursively, without integration.

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#### CHAPTER I

#### INTRODUCTION

The study of anharmonic oscillators is a subject of very general interest. The quartic anharmonic oscillator described by the Hamiltonian  $H = p^2 + x^2 + \lambda x^4$  (  $\lambda > 0$ anharmonicity constant) is the simplest example of nonlinear oscillation in classical mechanics (Krylov and Bogoliubov The investigation of the same problem in quantum 1943). mechanics is useful as a model problem in molecular physics and field theory In molecular vibrations, the potential functions are quite often of the mixed harmonic-quartic type. Indeed, in some cases of interest, due to accidental cancellation of the quadratic terms, the potential functions become nearly pure quartic (see Chan and Stelman 1963, Reid 1970). The quartic anharmonic oscillator is of particular interest in field theory because it is a model of  $\lambda \phi^4$  quantum field theory in one-dimensional space-time. The investigations from the field theory point of view are referred to in the work of Bender and Wu (1976). A general account on the anharmonic oscillator problem may be found in Hioe and Montroll (1975) and Hioe, MacMillen and Montroll (1976).

There have been persistent attempts for finding the eigenvalues of the quartic anharmonic oscillator since the beginning of quantum mechanics. In the earliest attempts the formal Rayleigh Schrödinger perturbation method was applied to solve this eigenvalue problem in which the  $\lambda x^4$ term was considered as the perturbation on the harmonic oscillator Hamiltonian  $H_0 = p^2 + x^2$ . Bender and Wu (1969) calculated nearly 70 expansion coefficients of the perturbation series in powers of  $\lambda$  for the ground state energy and observed an unexpected rapid increase in their magnitudes. A detailed analysis then revealed that the perturbation series for the eigenvalues in powers of  $\lambda$  is not convergent for any positive value of  $\lambda$  , no matter how Bender and Wu (1969) studied the analytic properties of the eigenvalues  $\mathbf{E}_{\mathbf{n}}(\pmb{\lambda})$  (n being the quantum number) in the complex  $\lambda$  plane and showed that there is a third order branch point at  $\lambda = 0$ . It was further observed that  $\lambda = 0$ is not an isolated singularity but is a limit point of an infinite number of branch points of  $E_n(\lambda)$ . Simon (1970) proved the above properties of  $E_n(\lambda)$  by more rigorous and technically sound arguments. In the language of perturbation theory the nonanalyticity of  $\mathbf{E}_{\mathbf{n}}(\lambda)$  implies that the perturbation series of  $\mathbb{E}_{n}(\lambda)$  in powers of  $\lambda$  is non-convergent.

The following scaling argument due to Symanzik is significant in this connection. On performing the scaling

transformation  $x \to ax$ ,  $p \to a^{-1}p$  on the Hamiltonian  $H(k, \lambda)$ =  $p^2 - kx^2 + \lambda x^4$  we obtain

$$H(k,\lambda) = \frac{1}{a^2}H(a^4k, a^6\lambda) . \qquad (1.1)$$

Since the above scaling transformation is implementable by unitary transformation the two sides of eqn. (1.1) have identical eigenvalues. The relation (1.1) thus ensures that the eigenvalue problem of the quartic anharmonic oscillator may be completely described in terms of the reduced Hamiltonian  $H(1,\lambda) = p^2 + \pi^2 + \lambda x^4$ . Setting  $a^6 = 1/\lambda$ , one obtains

$$E_{n}(1,\lambda) = \lambda^{1/3} E_{n}(\lambda^{-2/3},1)$$
 (1.2a)

Hence,

$$E_{n}(1,\lambda) \sim \lambda^{1/3} E_{n}(0,1) \text{ as } \lambda \rightarrow \infty$$
. (1.2b)

Since  $E_n(0,1)$  is independent of  $\lambda$ ,  $E_n(1,\lambda) \sim c_n \lambda^{1/3}$  for large  $\lambda$ .

The difficulties in perturbation approach may also be seen if one considers the equation  $H\psi = E\psi$  in momentum representation (Hioe and Nontroll 1975):

$$(\lambda \frac{d^4}{dp^4} - \frac{d^2}{dp^2} + p^2) \psi = \mathbb{E} \psi.$$
 (1.5)

The perturbation parameter  $\lambda$  appears here as the coefficient of the highest degivative. It is well known that the expansion of the solutions of such differential equation

in power series of the small parameter is non-convergent (VanDyke 1964). It is therefore not so surprising that the Rayleigh Schrödinger perturbation expansion of the anharmonic oscillator eigenvalues in powers of  $\lambda$  fails to con-The perturbation treatment of this problem in classical mechanics is instructive in this connection. consists of obtaining approximate solutions with the help of the expansion of the displacement in power series of the anharmonicity constant. It leads to solutions that contain secular terms like  $t^m$  sin  $\alpha t$ ,  $t^m$  cos  $\alpha t$  in which the time 't' appears outside the sine and cosine symbols. The secular terms in a finite term expansion introduce nonperiodic solutions and cause the calculated displacements to become arbitrarily large at large t. Further, the total energy becomes a function of time, violating the energy conservation principle (Bogoliubov and Mitropolsky 1961). The difficulties arising in the quantum mechanical perturbation expansion may be viewed in this context.

A detailed analysis of the perturbation series of  $E_n(\lambda)$  shows that it is asymptotic in nature (Simon 1970). Buch series are usually summed uniquely through various summability techniques such as Stieltjes Padė or Borel methods. Loeffel et al. (1969) have proved that the perturbation series sums under Padė approximation to the actual eigenvalue. The Padė approximation in general

consists of replacing the power series by a sequence of functions f(N.N) of the form of a polynomial of degree M divided by another polynomial of degree N. (1970) calculated the ground state energy by this approximation for various values of  $\lambda$ . His results show that the Padé approximants converge quickly for  $0 < \lambda < 1$  but for  $\lambda$  >1 the rate of convergence is not very good. et al. (1970) described how improved values of  $E_{\mathcal{C}}(\lambda)$  can be obtained by using Pade approximants to the Borel summability Reid (1967) showed that the perturbation series can also be summed by converting it into equivalent continued fraction and obtained the ground state as well as a few excited state eigenvalues for various values of  $\lambda$ . agreement of the eigenvalues obtained from these various summability methods with the corresponding accurate eigenvalues is found to be poor unless n and  $\lambda$  are sufficiently The scope of these methods is therefore limited to small n, small \u03bb values only.

Various variational and numerical methods have been widely employed by many authors either to the quartic anharmonic oscillator problem or to the associated problem of the pure quartic oscillator (H =  $p^2 + \lambda x^4, \lambda > 0$ ). Calculations are generally done in the harmonic oscillator basis { $x^m e^{-x^2/2}$ } and the usual technique is to truncate and diagonalize a large but finite matrix. Results of such

calculations are summarised in Table (I.1). They differ from one another either in the method of diagonalization or in the size of the basis used. Some calculations for upper and lower bounds of first few eigenvalues with varying anharmonicity were done by Bazley and Fox (1961) and Reid (1965). The procedure for lower bounds used by Bazley and Fox was to construct intermediate Hamiltonians  $H^k$  such that  $H^0 < H^1 < H^2$  ... <H and to determine eigenvalues of successive  $H^k$ . Reid used the method of Löwdin (1965) for obtaining the lower bounds. The upper bounds were calculated by employing the usual Rayleigh-Ritz variational approach. It may be noted that in these calculations the gap between the upper and the lower bounds increases rapidly on increasing n or  $\lambda$  (e.g. in Reid's work the gap, which is  $O(10^{-15})$  for  $E_C(\lambda=0.25)$ , becomes  $O(10^{-1})$  for  $E_C(\lambda=1.0)$ ).

Biswas et al. (1973) used the 'Hill determinant' method to obtain first eight eigenvalues of the quartic anharmonic oscillator for values of  $\lambda$  in the range  $0 < \lambda \le 100$ . They used an expansion in terms of the functions  $\{x^m e^{-x^2/2}\}$  for the eigenfunctions and obtained eigenvalues by finding the roots of a sequence of characteristic polynomials in E. The polynomials corresponded to various order truncations of the Hill determinant. For higher eigenvalues or for higher  $\lambda$  the numerical errors in their work become too severe

TABLE (I.1); Comparison	arison	of the Resu	Jo	ons	Veriational	and Numerical Methods.
Authors	Year	System	No. or basis functions used	n	Control of the Contro	Accuracy of $\vec{\mathrm{E}}_{\mathrm{n}}(\lambda)$ and other remarks
McNeeny and Coulson	1948	Quartic Oscillator	ω	0 to 4	1.0	$7$ figures for $\overline{\mathrm{E}}_{\mathrm{O}^{s}}$ 5 figures for $\overline{\mathrm{E}}_{\mathrm{Q}}$
Chan and Stelman	1963	-	50	0 to 19	1.0	8 significant figures (for n $\ge 17$ E <sub>n</sub> ( $\lambda$ ) are inaccurate in last 5-4 figures)
V <b>e</b> scelius and Neff	1968	-	ì	0 to 19	1.0	6 to 8 significant figures (the method uses continued fraction technique)
Bell et al.	1970	- -	800	0 to 49	J. C	7 to 9 significant figures (for n > 17 $\rm E_{\rm n}(\lambda$ ) are inaccurate in last figures)
Reid	1970	San- Care	100	C to 23	1.0	12 significent figures
Bazley and Fox	1961	Quartic Anhar- monic oscil- lator	۲. ت	0,2,4,	0.1 40 1.0	5-6 significant figures for ${\tt E}_0$ , none for ${\tt i}_8$
Chan, Stelman and Thompson	1964	Ξ	20	0 to 9	i	4-5 significant figures for lower eigenvalues, 1-2 significant figures for higher eigenvalues.
Re <u>i</u> d	1965	<u>:</u>	20	0 to 3	0,1 to 1,0	15 significant figures for E <sub>0</sub> ( $\lambda$ = 0.25), 2 significant figures for E <sub>9</sub> ( $\lambda$ = 1.0)
Biswas et al.	1973	-		0 to 7	0,1 to 10C	14 significant figures for 50 6-9 significant figures for n > 1.

The results of the variational and numerical calculations show that these methods are suitable for the evaluation of a few lower eigenvalues and for moderate anharmonicities. The accuracy of the results is seriously affected on increasing n or  $\lambda$  and is not necessarily improved by increasing the size of the basis. Further, the eigenfunctions obtained from variational calculations are far less accurate than the corresponding eigenvalues. Evaluation of matrix elements of operators using such eigenfunctions is not expected to yield values of known or definite accuracy.

The WKB method has been used for obtaining approximate eigenvalues for high n. In the WKB approximation (Titchmarsh 1961),

$$E_n(0,1) = C(n + \frac{1}{2})^{4/3}, C = 2^{2/3}(1.376)$$
 (1.4)

It is known from eqn. (1.2b) that the quartic anharmonic oscillator eigenvalues  $E_n(1,\lambda) \simeq \lambda^{1/3} E_n(0,1)$  in the large  $\lambda$  limit. Hence, for large n, large  $\lambda$  -

$$E_n(1,\lambda) \simeq C \lambda^{1/3} (n + \frac{1}{2})^{4/3}$$
 (1.5)

A more sophisticated WKB expression for  $E_n(1,\lambda)$  is obtained by Hioe and Montroll (1975); see eqn. (1.6.c) later.

The most comprehensive work on the anharmonic oscillators is due to Hioe and Montroll (1975) and Hioe et al. (1976) They distinguished two limiting regimes of values of n and  $\lambda$ .

In one regime the energy eigenvalues differ slightly from the harmonic oscillator levels; in the other they differ slightly from the pure quartic oscillator eigenvalues. These regimes are called the 'near harmonic' regime and the 'near quartic' regime, respectively. Between these two regimes lies the 'boundary layer' in which the eigenvalues are neither 'near harmonic' nor 'near quartic'. 'They developed fast converging algorithms for computing the eigenvalues in small n regime by writing the eigenvalue problem in Bargmann representation and solving the associated difference The eigenvalues were thus computed to 8-9 significant figures for n = 0, l, ... 8 and for values of  $\lambda$  in the range  $.004 \le \lambda \le 40000$ . They also constructed several simple formulae for  $E_n(\lambda)$  with different ranges of validity which when combined give good approximations (about 8 significant figures) to  $\textbf{E}_{n}(\lambda\,)$  except in the 'boundary layer'. The formulae with their ranges of validity are as follows:

(a) For the near harmonic regime

$$E_{n}(\lambda) = (2n+1) + \frac{3}{4}\lambda \left\{ 1 + 2n(n+1) \right\} - \lambda^{2} \left\{ \frac{(n+1)(n+\frac{3}{2})^{2}(n+2)}{[4+3\lambda(2n+3)]} + \frac{(n+1)(n+2)(n+3)(n+4)}{32[4+3\lambda(2n+5)]} - \frac{n(n-\frac{1}{2})^{2}(n-1)}{[4+3\lambda(2n-1)]} - \frac{n(n-1)(n-2)(n-3)}{32[4+3\lambda(2n-3)]} \right\} + 0 (\lambda^{3})$$
(1.6a)

(b) For small n, large  $\lambda$  region

$$\bar{\mathbf{E}}_{n}(\lambda) = \lambda^{1/5} \left[ \varepsilon_{n} + \alpha_{n} \lambda^{-2/3} + \beta_{n} \lambda^{-4/5} + \ldots \right], \quad (1.6b)$$

where the constants  $\epsilon_n$ ,  $\alpha_n$  and  $\beta_n$  were determined by fitting eqn. (l.6b) to the numerical values of  $E_n(\lambda)$  for each state up to n=10.

(c) For large n, large  $\lambda$  region

$$E_{n}(\lambda) = \lambda^{1/3} \left[ c \left\{ \left( n + \frac{1}{2} \right) + \frac{\delta}{\left( n + \frac{1}{2} \right)} \right\}^{4/3} + a \left( n + \frac{1}{2} \right)^{2/3} \lambda^{-2/3} + b \lambda^{-4/3} + \ldots \right].$$
(1.0c)

The above expression results from a detailed investigation of the WKB approximation formula and consists of expanding the elliptic integrals in the WKB formula in a series in  $E_n$ ; the constants c, a and b are identified from this expansion. It may be mentioned here that Mathews and Usuaran (1972) also obtained some approximate formulae through a semiclassical treatment

Recently, hawhers and Covindarajan (1977) used a 'residue squaring method' for the iterative diagonalization of the quartic anharmonic oscillator Hamiltonian in which the  $\lambda x^4$  term is assumed to be a perturbation on the rest. In this work the off diagonal part (of order  $\lambda$  compared to the diagonal part) is successively reduced to orders  $\lambda^2$ ,  $\lambda^4$ ,  $\lambda^8$ , .... They obtained four lowest even parity eigenvalues and an approximate analytic formula for  $E_n(\lambda)$ , similar to the

equation (1.6a), which gives good approximations to the eigenvalues in the near harmonic regime.

It is seen from the above discussion that the various methods, which have been applied to solve the anharmonic oscillator eigenvalue problem, are suitable only in some particular regime of  $(n,\lambda)$  and do not give eigenvalues to the same accuracy when extended to other regimes. Moreover, then do not yield accurate eigenfunctions. In contrast, we use a method (Banerjee 1976) to obtain the eigenvalues of the quartic anharmonic oscillator which applies with uniform and arbitrarily high accuracy for all values of n and The method also yields eigenfunctions of accuracy comparable with that of the eigenvalues which are used for the computation of high accuracy matrix elements. In the next chapter the method is described and the eigenvalues and the eigenfunctions of the quartic anharmonic oscillator and the associated problem of the pure quartic oscillator are obtained. The eigenvalues, accurate to 15 significant figures, are presented for various values of  $(n,\lambda)$  covering all the different regimes. We show in Chapter III that the method may be extended to solve general anharmonic oscillator  $(H = p^2 + x^2 + \lambda x^{2\mu}, \mu = 3,4,...)$  eigenvalue problem. Eigenvalues of the sextic ( $\mu$ =3) and the octic ( $\mu$ =4) anharmonic oscillators are thus obtained accurate to 15 significant figures in all regimes of  $(n,\lambda)$ . In Chapter IV the eigenvalue spectrum of the double minimum oscillator ( $H = p^2 - x^2 + \lambda x^4$ ,  $\lambda > 0$ ) is investigated. The lower eigenvalues of the double minimum oscillator are closely bunched in pairs for small  $\lambda$ . These eigenvalues have been evaluated accurately using the same method. A WKB expression for the splitting between the eigenvalues bunched in pairs is obtained and the WKB values are compared with the corresponding accurate values. In Chapter V the transition moments between the anharmonic oscillator energy eigenstates are obtained from the computed eigenvalues and eigenfunctions. Further, the multipole transition moments are shown to satisfy an exact linear recurrence relation which is valid for any polynomial potential.

#### CHAPTER II

### THE QUARTIC ANHARMONIC OSCILLATOR

In this chapter we determine accurate eigenvalues and eigenfunctions of the quartic anharmonic oscillator  $(H=p^2+x^2+\lambda x^4,\,\lambda>0,\,p=-i\,\frac{d}{dx})$  for various values of the quantum number n and the anharmonicity constant  $\lambda$ . The existence of two distinct regimes of values of  $(n,\lambda)$ , namely the 'near harmonic' and the 'near quartic', separated by a 'boundary layer' implies different oscillation properties of the corresponding eigenfunctions. We assert that this fact must be explicitly included in solving the eigenvalue problem. Following Banerjee (1976), it is shown in the next section that the basis functions (used in the solution of the eigenvalue problem) may be appropriately scaled to simulate the oscillation properties of the eigenfunctions in all regimes.

## II.l Scaling and the Appropriate Scaling Formula

The eigenfunctions  $\psi_n(x; \lambda)$  may be expanded in the basis  $\{x^m e^{-\alpha x^2}\}_{as}$ :

$$\Psi_{n}(x, \lambda) = e^{-\alpha x^{2}} \sum_{m=0}^{\infty} a_{m} x^{m}, \qquad (2.1)$$

where the scaling is introduced through the parameter 'a'. For an effective expansion the scaling a is chosen such that a sufficient number of the lover members of the basis functions (at least n for the n-th state) have their main contribution in the region of oscillation and outside it they decay Since the region of oscillation depends on monotonically n and  $\lambda$ , the scaling  $\alpha$  also depends on n and  $\lambda$ . A simple criterion for the appropriate scaling is obtained as follows: The region of oscillation for the n-th eigenfunction of the quartic anharmonic oscillator (for sufficiently large n)  $\sim (E_n/\lambda)^{1/4}$ . In the WKB approximation  $E_n(\lambda) \simeq C\lambda^{1/3}(n+\frac{1}{2})^{4/3}$ . Therefore the region of oscillation  $\sqrt{1/6}(n+\frac{1}{2})^{1/3}$ . The exact n-th eigenfunction has n zeros in the region of oscil-Hence setting the region of oscillation for the n-th eigenfunction equal to the width of the n-th basis function  $(\sqrt[\alpha]{(n/\alpha)})$  we obtain

$$\alpha(n, \lambda) \sim (n + \frac{1}{2})^{1/3} \lambda^{1/3}$$
.

This puts the span of the first n (or a number proportional to n) basis functions in the region of oscillation of the n-th eigenfunction for all n and  $\lambda$ , just as required for an effective expansion. In view of the WKB estimate and the large  $\lambda$  assumption implicit in the derivation of the above scaling formula, it is not expected to be good when n and or  $\lambda$  is small. However, for  $\lambda \to 0$  or for small n the

scaling must approach the value 1/2 appropriate for the harmonic oscillator. Hence the scaling formula for all regimes of  $(n,\lambda)$  is

$$\alpha(n,\lambda) = \frac{1}{2} + (n + \frac{1}{2})^{1/3} \lambda^{1/3}$$
 (2.2)

The effect of using an appropriately scaled basis is remarkable. It is now possible to compute the eigenvalues in any regime of  $(n,\lambda)$  with arbitrarily high accuracy.

It is to be noted that the different regimes of  $(n,\lambda)$  are distinguished according to the above scaling formula. Thus

$$(n + \frac{1}{2})^{1/3} \lambda^{1/3} \ll \frac{1}{2}$$
 is the near harmonic regime,  $(n + \frac{1}{2})^{1/5} \lambda^{1/3} \gg \frac{1}{2}$  is the pure quartic regime,  $(n + \frac{1}{2})^{1/3} \lambda^{1/3} \simeq \frac{1}{2}$  is the boundary layer between the above two regimes.

The value of the combination  $(n+\frac{1}{2})\lambda$  is seen to determine the regime to which an eigenvalue belongs. The importance of the above combination of n and  $\lambda$  in determining the various regimes was also recognised by Hioe et al. (1976) on essentially empirical grounds. It is in this work that the combination  $(n+\frac{1}{2})\lambda$  is shown to determine the characteristic scaling in a given regime of  $(n,\lambda)$  through the relation (2.2). This observation leads to the construction of a scale adapted

basis and makes possible a uniform treatment of the problem in all regimes of  $(n,\lambda)$ .

### II.2 Method

The method applied for obtaining the eigenvalues is described in this section. The Schrödinger equation for the quartic anharmonic oscillator is

$$\left[-\frac{\mathrm{d}^2}{\mathrm{d}x^2} + x^2 + \lambda x^4\right] \psi(x;\lambda) = \mathbb{E}(\lambda) \psi(x;\lambda), \qquad (2.3)$$

where the eigenfunctions  $\psi_n(x;\lambda) \to 0$  as  $x \to \pm \infty$ . The expansion (2.1) on substitution into the above equation yields the following 4-term linear recurrence relation, connecting the alternate expansion coefficients  $\{a_m\}$ ,

$$(m+1)(m+2) a_{m+2} + (E-4\alpha m-2\alpha) a_m + (4\alpha^2-1) a_{m-2} - \lambda a_{m-4} = 0$$

$$(2.4)$$

the above recurrence relation may be rewritten in the following notations:

$$a_{m+2} + d_{m,m}a_{m} + d_{m,m-2}a_{m-2} + d_{m,m-4}a_{m-4} = 0$$
, where

$$d_{m,m} = \frac{(E - 4\alpha m - 2\alpha)}{(m+1)(m+2)}, d_{m,m-2} = \frac{(4\alpha^2 - 1)}{(m+1)(m+2)},$$

$$d_{m,m-4} = -\frac{\lambda}{(m+1)(m+2)}.$$
(2.5)

Since the Hamiltonian for the system has even symmetry, the solutions of the Schrödinger equation are either even or odd functions of x. The even and odd parity solutions are obtained respectively by assigning the initial conditions (i)  $a_0=1$ ,  $a_1=0$  (ii)  $a_0=0$ ,  $a_1=1$ . The recursion(2.5) may be viewed as an infinite set of linear homogeneous equations in the unknowns  $\{a_m\}$ . For the self consistency the infinite determinant  $\mathbf{A}(\mathbf{E})$  formed from the coefficients of  $\{a_m\}$  must vanish. It gives

The roots of the above transcendental equation are the eigenvalues. This mode of writing the characteristic equation in the form of an infinite determinant is well known from the eigenvalue problem associated with Hill's equation (Whittaker and Watson 1927). Denoting the truncated determinant formed by omitting all rows and columns beyond the element  $d_{m,m}$  as  $\Delta_{m+2}(E)$ , it may be noted that  $\Delta_m(E)$  is a polynomial in E

$$\Delta_{m+2}(E) - d_{m,m}(E) \Delta_{m}(E) + d_{m,m-2} \Delta_{m-2}(E) - d_{m,m-4} \Delta_{m-4}(E) = 0,$$
(2.7)

obtained by expanding the determinant  $\Delta_{m+2}(E)$ . The value of the determinants  $\Delta_m(E)$  may thus be computed successively upto any order in terms of  $\Delta_0(\text{or }\Delta_1)$  using (2.7). The recurrence relation (2.7) on differentiation with respect to E yields

$$\Delta_{m+2}^{i}(E) - d_{m,m}^{i} \Delta_{m}(E) - d_{m,m}(E) \Delta_{m}^{i}(E) + d_{m,m-2} \Delta_{m-2}^{i}(E) - d_{m,m-4} \Delta_{m-4}^{i}(E) = 0, \qquad (2.8)$$

from which  $\Delta_m^{'}(\Xi)$  may be computed upto any order recursively. The recursions (2.7) and (2.8) are numerically stable. The initial estimates for the eigenvalues required in the Newton's method may be obtained for low n by evaluating a

sufficiently large order determinant  $\Delta_{\mathbb{M}}(E)$  from the recursion (2.7) at various E points. Opposite signs of  $\Delta_{\mathbb{M}}(E)$  for two neighbouring E values indicate—that an eigenvalue is crossed which provides sufficiently accurate estimate for the Newton's method. For high n (and not too low  $\lambda$ ) the corresponding WKB approximations of the eigenvalues are good initial estimates. The procedure for obtaining initial estimates in the (high n, low  $\lambda$ ) region of the 'boundary layer' is described in Appendix A.

The actual computation of the eigenvalues may now be performed in the following manner. An initial estimate  $(\Xi=\Xi_{\rm initial})$  is fed into the recursions (2.7) and (2.8) containing the appropriate value for  $\alpha$ . The recursions are then continued on a computer until the corrections, given by the Newton's formula

$$\delta E(m) = -\{ \Delta_m(E)/\Delta_m'(E)' \}_{E=E_{\text{initial}}}, \qquad (2.9)$$

stabilize to a prescribed extent (see Wilkinson 1965). The corrected value for E is then fed back in the second step as the initial value. This is continued till the required accuracy in the computed eigenvalues is reached. Due to the quadratic convergence of Newton's method it is possible to refine a rather crude initial estimate for an eigenvalue (say, within a few percent) to a 15 figure accuracy in 4 or 5 steps for all eigenvalues. Some typical examples to

elucidate this are given in Table (II.1). In principle, the method can be carried to an arbitrary high accuracy. The accuracy of the computation is limited only by the precision of the arithmetic used (16 significant figures in IBM 7044).

### II.3 Eigenvalues

Very accurate eigenvalues of the quartic anharmonic oscillator and the associated problem of the pure quartic oscillator (H =  $p^2 + \lambda x^4, \lambda > 0$ ) have been obtained using the method described in the previous section and the scaling formula

$$\alpha(n,\lambda) = \frac{1}{2} + (1.2 \text{ to } 1.4) (n + \frac{1}{2})^{1/3} \lambda^{1/3}$$
. (2.10)

The constant within the bracket (1.2 to 1.4) has been set empirically by finding the values of the scaling  $\alpha$  for which the computed eigenvalues stabilize the earliest. The constant (1.2 to 1.4) is found to work admirably in the entire range of  $(n,\lambda)$ . The calculation of the eigenvalues for the pure quartic oscillator is similar to that done for the quartic anharmonic oscillator. The only difference is in the value of  $d_{m,m-2}$  in equations (2.5) to (2.8), which in the pure quartic oscillator case is equal to  $(4\alpha^2/(m+1)(m+2))$ . The computations were done on IBM 7044 computer using double precision arithmetic (16 digits mantissa). The eigenvalues were evaluated to

<sup>\*</sup> Tables referred to in any chapter are given at the end of that chapter.

16 significant figures and then rounded off to 15 figures for the Tables.

The eigenvalues of the quartic anharmonic and the pure quartic oscillator for various values of  $(n, \lambda)$  covering all regimes are listed in Tables (II.2 to II.5). In Table (II.2) we compare our results of  $E_n(\lambda=1)$  for n=0, 10, 100, 1000 and 10000 with the corresponding results of various earlier calculations. In Table (II.3) the first 50 eigenvalues of the quartic anharmonic and the pure quartic oscillators are given for  $\lambda=1$ . The eigenvalues for any other value of  $\lambda$  in the case of pure quartic oscillator can be obtained from the corresponding values for  $\lambda = 1$  through the exact scaling relation  $E_n(\lambda)$ =  $\lambda^{1/3}E_n(1)$ . For the quartic anharmonic oscillator different eigenvalues for various values of  $\lambda$  between .00001 and 40000 were computed, and are presented in the Table (II.4). includes the eigenvalues for the (high n, low  $\lambda$ ) region of the 'boundary layer'. The computation of eigenvalues in this region of the boundary layer is found to be the most difficult in the earlier literature. None of the approximation formulae (1.6a,b,c) constructed by Hioe et al. (1975) are adequate for this (high n, low  $\lambda$ ) region. In the present work the eigenvalues in this region (e.g. eigenvalues corresponding to n=100,  $\lambda = 10^{-4}$  and n=1000,  $\lambda = 10^{-3}$  in Table (II.4)) are obtained by the same technique and with the same accuracy as any other,

highlighting the scope of computations with an appropriately scaled basis. In Table (II.5) we focus on the regimes of extreme values of  $(n,\lambda)$ . Hioe et al. (1975, 1976) give different formulations of the eigenvalue problem in each of these regimes which cannot be extended into other regimes because of the boundary layer in between. In contrast, we have obtained eigenvalues in each of these regimes by the same formulation and with the same accuracy.

# II.4 Stability of Zeros of $\Delta_{m}(E)$

The recursive evaluation of the determinants  $\Delta_m(E)$  and the stability of their zeros will now be considered in some detail. It may be noted that the recursion (2.7) is obtainable from the recursion (2.5) by replacing  $a_m$  with  $\Delta_m$  and changing the sign of every alternate term. This prescription is valid when the coefficient of the highest order term in the  $\{a_m\}$  recursion is set unity (by properly dividing, if necessary). Then

$$\Delta_{\rm m} = (-1)^{\rm m/2} a_{\rm m}, \, m = 0,2,4,...,$$

$$\Delta_{\rm m} = (-1)^{(\rm m-1)/2} a_{\rm m}, \, m=1,3,5,...$$
(2.11)

In the case when the coefficient of the highest order term is not unity, the  $\{a_m\}$  recursion is

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$$\Delta_{\rm m} = (-1)^{{\rm m}/2} a_{\rm m}, m = 0,2,4,...,$$

$$\Delta_{\rm m} = (-1)^{({\rm m}-1)/2} a_{\rm m}, m=1,3,5,... \qquad (2.11)$$

In the case when the coefficient of the highest order term is not unity, the  $\{a_m\}$  recursion is

$$d_{m,m+2} a_{m+2} + d_{m,m} a_m + d_{m,m-2} a_{m-2} + d_{m,m-4} a_{m-4} = 0$$
. (2.12)

The corresponding infinite determinant formed in this case is

The truncated determinants  $\overline{\Delta}_{m+2}(E)$  formed by omitting all rows and columns beyond the element  $d_{m,m}$  in  $\overline{\Delta}(E)$  may be expanded in terms of the truncated determinants of lower orders. It provides the following 4-term recurrence relation

$$\bar{\Delta}_{m+2}(\Xi) - d_{m,m} \bar{\Delta}_{m}(E) + d_{m-2,m}d_{m,m-2} \bar{\Delta}_{m-2}(E)$$

$$- d_{m-4,m-2}d_{m-2,m}d_{m,m-4} \bar{\Delta}_{m-4}(E) = 0 . \qquad (2.14)$$

The determinants  $\{\overline{\Delta}_{\mathrm{m}}(\mathtt{E})\}$  are related to  $\{\Delta_{\mathrm{m}}(\mathtt{E})\}$  by

$$\bar{\Delta}_{m}(E) = d_{m-2,m} d_{m-4,m-2} \dots d_{02} \Delta_{m}(E), m=0,2,4,\dots,$$

$$\bar{\Delta}_{m}(E) = d_{m-2,m} d_{m-4,m-2} \dots d_{13} \Delta_{m}(E), m=1,3,5,\dots.$$
(2.15)

1

The asymptotic behaviour of the solution of the Schrödinger equation  $(2.3) \sim \exp(\pm |x|^3/3)$ . It requires

$$a_{m+2}/a_{m-4} \sim \lambda/m^2$$
,  $m \to \infty$ , (2.16)

in the series solution (2.1). It follows immediately from the relation (2.11) that for the determinants  $\{\Delta_m\}$  it is required that

$$\Delta_{m+2}/\Delta_{m-4} \sim -\lambda/m^2 , \qquad m \to \infty . \qquad (2.17)$$

We observe during the recursive evaluation of the value of  $\Delta_{m+2}(\mathbb{E})$  that for sufficiently large m the second and third terms in the recursion (2.7) become order of magnitudes smaller than the last term and the asymptotic dependence (2.17) is satisfied. The asymptotic relation (2.17) implies a sequence of decreasing determinants beyond a sufficiently This zeroing for large m must be isolated from the determination of the eigenvalues which are the zeros of  $\Delta$  (E) for values of E. It is possible to achieve this quite simply by redefining the determinants  $\Delta_m(E)$  after multiplying them with a large number whenever, while applying Newton's method, the recursively computed determinants become too This renormalization amounts to starting small in magnitude. the recursion with a higher value of the arbitrary constant  $\Delta_{\cap}$  (or  $\Delta_{1}$ ). More generally, other recursively connected sequences of determinants  $\{\widetilde{\Delta}_{m}(E)\}$  may be defined such that

the zeros of  $\tilde{\Lambda}_m(E)$  and  $\Lambda_m(E)$  are compon but  $\tilde{\Lambda}_m(E)$  may be given any desired asymptotic behaviour for large m. This is done by multiplying the recursion (2.5) by a function of m say f(m). The corresponding infinite determinant  $\tilde{\Lambda}(E)$  and its various order truncations are related to the respective quantities for  $\Lambda(E)$  by the relation

$$\tilde{\Delta}_{m}(E) = f(m-2) f(m-4) \dots f(\cup \text{ or } 1) \Delta_{m}(E), (2.18)$$

where the the contains f(0) (or f(1)) for the even (or odd) eigenvalues. Clearly  $\tilde{\Delta}_m(E)$  can be given any asymptotic behaviour for large m by properly choosing the function f(m). Since f(m) is independent of E by definition, the zeros of  $\tilde{\Delta}_m(E)$  and  $\Delta_m(E)$  are common and they are equally suited for the computation of the eigenvalues. The 'renormalization' suggested above is a special case of this multiplication in which all rows are left intact except one which is multiplied by a large number.

To see the stability of the zeros of  $\Delta_m(E)$  as  $m \to \infty$  we first consider the same problem for the harmonic oscillator ( $H = p^2 + x^2$ ) which is exactly soluble. In the case of the harmonic oscillator the zeros of the characteristic polynomial  $\Delta_m(E)$  are real and the ratio of successive polynomials is

$$\frac{\Delta_{m+2}(E)}{\Delta_{m}(E)} = -\frac{(2m+1-E)}{(m+1)(m+2)}.$$

The zeros of  $^{\Delta}_{m+2}(E)$  consist of all zeros of  $^{\Delta}_{m}(E)$  plus a zero at E=2m+1. Thus the eigenvalues resulting from the solution of  $^{\Delta}_{m}(E)=0$  are reproduced exactly by the solutions of  $^{\Delta}_{m+2}(E)=0$  for all m. This is characteristic of an exactly soluble problem. For the quartic anharmonic or the pure quartic oscillator the zeros of successive order polynomials are different. For a given E and sufficiently large m, however,

$$\frac{\Delta_{m+2}(E)}{\Delta_{m-4}(E)} - \frac{\lambda}{m^2}, \quad m \to \infty .$$

## II.5 Checks for the Eigenvalues

The confidence in the accuracy of the computed eigenvalues is derived from the rollowing checks:

(i) Computations were done with several different initial estimates for each eigenvalue. The intermediate numbers involved in the computation are different for different initial estimates but the final results for the eigenvalues remain the same in all cases.

- (ii) The eigenvalues were computed using several values of the scaling  $\alpha$  in the range given by eqn. (2.10). Its effect is that the stabilization of an initial estimate for an eigenvalue to 10 figures occurs at slightly different point in recursions (2.7) and (2.8). Each  $E_n(\lambda)$  obtained in this work was thus checked and confirmed by using 5 or 4 different values of  $\alpha$ .
- (iii) Three separate computations using an increasing, a decreasing and a nearly flat sequence of determinants (see section II.4) yielded the same values for  $\mathbb{F}_n(\lambda)$ .
- (iv) Sufficiently large order determinant  $\Delta_{\rm M}$  containing the appropriate scaling is evaluated for two neighbouring values of E. Opposite signs of  $\Delta_{\rm M}$  indicates that an eigenvalue is crossed. The computed eigenvalues are thus tested and in the process are upper and lower bounded in the last significant figure.

# II.6 Eigenfunctions

When E is set equal to a computed eigenvalue in recursion (2.5) the resulting coefficients  $\{a_m(E)\}$  provide a very convenient representation for the corresponding eigenfunction through the expansion (2.1). The following properties of the solutions of the Schrödinger equation  $\psi(x,E)$  are important to note in this connection (Titchmarsh 1961):

- (i) The eigenfunction associated with the n-th eigenvalue  $\psi_n(x;E)$  has n zeros in the classical region and  $\psi_n(x;E_n) + 0$  as well as  $\psi_n'(x;E_n) + 0$  as  $x + \pm \infty$ .
- (ii) For E not exactly equal to an eigenvalue,  $\psi$  (x;E) can have at most one zero in the nonclassical region on both sides of the axis and as  $x \to \pm \infty$  it goes either to  $\pm \infty$  or  $\pm \infty$ .
- (iii) The blow up of  $\psi(x;E)$  starts in the nonclassical region and shifts to larger |x| as E approaches an eigenvalue.
- (iv) As E crosses an eigenvalue,  $\psi(x;E)$  changes its sign of blow up in the nonclassical region and tends to infinity with opposite sign.

The expansion coefficients  $\{a_m(E)\}$  are evaluated recursively from (2.5) for various computed eigenvalues. It is observed that the sign of  $\{a_m\}$  stabilizes after a certain sufficiently large index either to plus or minus eight which implies that the computed  $\psi(x,E) + +\infty$  or  $-\infty$  as  $x \to \pm \infty$ . However, this large |x| behaviour does not affect the computation of the accurate eigenfunctions significantly. Since the eigenvalues used are accurate to 15 significant figures, the computed eigenfunctions reach extremely small values in the nonclassical region before the blow up starts. The part of the computed eigenfunctions where the blow up occurs for large |x| may therefore be replaced by zero without losing much information.

To test how well the computed eigenfunctions satisfy the Schrödinger equation we compare the two sides of  $H\psi(x)/\psi(x) = E$  at various points x. For the first ten eigenfunctions which we have computed for  $\lambda = 1$  the test equality is satisfied to 19-14 significant figures from x=0 to points well outside the classical region. For example, in the case of the 10th eigenfunction of the quartic anharmonic oscillator with  $\lambda=1$ , the test equality  $H\Psi_{10}/\Psi_{10}=\Xi_{10}$  is satisfied to at least 13 significant figures in the entire classical region (x  $\leq E_{10}^{1/4}$ ). At a point x = 1.5  $E_{10}^{1/4}$  the test equality is still satisfied to 10 significant figures, where the value of the computed eigenfunction  $\psi_{10}$  (x = 1.5  $E_{10}^{1/4}$ ) is  $\zeta(10^{-16})$  relative to  $\psi_{10}(x=0)=1$ . The accuracy of the computed eigenfunctions are also checked by evaluating them For two neighbouring values of E which upper and lower bound the eigenvalues in the 15th significant figure. Although, the sign of the expansion coefficients  $\{a_m\}$  stabilizes to all plus or all minus beyond a sufficiently high index m=M, the difference in the corresponding computed values of  $\psi(x;E)$ for these neighbouring values of E is found to be less than  $\text{O(10}^{-15})$  for all  $|\mathbf{x}| < \mathbf{x}_{A}$ , where  $\mathbf{x}_{A}$  is the distance from the origin to the point in the nonclassical region at which  $\psi(\mathrm{x};\mathrm{E})$  begins to increase in magnitude. Satisfying the virial theorem by the computed eigenfunctions was used in the earlier literature (e.g. Chan and Stelman 1963) to test t accuracy. However, the fulfilment of the virial theorem is a necessary but not sufficient requirement (Löwdin 1959).

The norm of the computed eigenfunction is,

$$\int_{-\infty}^{+\infty} |\psi_{n}(x)|^{2} dx = \sum_{m} \sum_{l} a_{m}^{(n)} a_{l}^{(n)} \int_{-x_{A}}^{+x_{A}} x^{m+l} e^{-2\alpha x^{2}} dx , \quad (2.19)$$

where the range of integration is truncted at  $x_A$  — the point in the nonclassical region at which the computed eigenfunction  $\psi_n(x)$  begins to increase in magnitude. The value of  $\psi_n(x)$  is sufficiently small for  $|x| > x_A$  as discussed above and the contribution to the normalization from the rest of configuration space is estimated to be  $<0(10^{-16})$ . The integrals in eqn. (2.19) are obtained recursively starting from the incomplete Gaussian integral  $\int\limits_0^1 e^{-\beta x^2} dx$  (see Appendix B). The plots of some normalized computed eigenfunctions are shown in Fig. (II.1).

## II.7 Features of the Method

In this section some features of the method are seen in comparison with the other methods used for this eigenvalue problem.

(i) It may be noted that no integration or diagonalization is necessary in this method which makes it attractive for the eigenvalue problems of the linear operators.

<sup>\*</sup> The vertical lines in the figure correspond to the classical turning points.

- (ii) In perturbation theory with  $\lambda$  as the small parameter, this problem belongs to the singular perturbation class. The uniform applicability of this method for all  $\lambda$  underlines its nonperturbative character.
- (iii) In view of the Section II.1 an expansion for the eigenfunctions like (2.1) with  $\alpha=$  constant may be called as a 'fixed scale' expansion. Such an expansion is suitable only in a small retime of values of  $(n,\lambda)$  where the scale happens to be close to the appropriate value and it becomes unfavourable in the other regimes of  $(n,\lambda)$ . The variational and most of the numerical methods applied earlier use the expansion

$$\psi_{\rm n} (x; \lambda) = e^{-x^2/2} \sum_{\rm m=0}^{\infty} a_{\rm m} x^{\rm m}$$
 (2.20)

in all regimes of  $(n, \lambda)$ . This expansion has a fixed scale  $\alpha = \frac{1}{2}$  and is suitable only in the 'near harmonic' regime. It is therefore not surprising that for higher n or  $\lambda$   $(n\lambda >> \frac{1}{8})$  the eigenvalues could not be accurately calculated in the above works.

(iv) The need for introducing a scaled basis was also realised by Reid (1970) who used the linear variation method for the pure quartic oscillator eigenvalue problem. However, in a variational framework the use of a scaled basis becomes intractably laborious for the following reasons. In a variational computation the first n (say) eigenvalues are

obtained together. Since the appropriate scaling is different for different n and  $\lambda$  a single scaling is not suitable for the computation of all n eigenvalues. A compromise scaling must therefore be used. But as n increases this compromise scaling becomes unfavourable for more and more eigenvalues. The wayout is to compute each eigenvalue separately using an appropriately scaled set of basis functions. This is intractably laborious in a variational scheme. In our method the use of an appropriately scaled basis merely requires that a proper value of the scaling  $\alpha$  obtained from formula (2.10) be used in recursions (2.5), (2.7) and (2.8). Since each eigenvalue is computed individually there are no carry over errors.

(v) Computation with a larger basis is very simply done in this method by continuing the recursions (2.5), (2.7) and (2.8) for increasing m. In contrast a variational calculation with a larger basis requires integration and the subsequent diagonalization of a large matrix which beyond a size is intractable. For instance, the 10000th eigenvalue of the quartic anharmonic oscillator stabilizes to a 15 figure accuracy (in 3 minutes on IBN 7044) at a point in the recursions which corresponds to the use of nearly 17500 terms in the expansion (2.1). A variational calculation of this size is inconceivable.

The method of infinite determinant for the eigenvalue (vi) problems (Whittaker and Watson 1927) was used by Kerner (1951) and Biswas et al. (1973). These attempts had a limited success. The primary reason for this is the use of a fixed scale expansion as discussed above. In the typical case of the quartic anharmonic oscillator (Biswas et al. 1973), only the lowest 8 eigenvalues could be obtained (using the expansion in the basis functions  $\{x^m e^{-x^2/2}\}$ ), until the numerical errors become too severe. Bosides, in this work, the procedure used for evaluating the eigenvalues consists of expanding the characteristic polynomials in powers of E and then finding its zeros. This is numerically inadvisable (see Fox and Mayers 1968) because the uncertainties in the coefficients of the polynomials are highly correlated and the expanded polynomials with rounded coefficients are badly conditioned with respect to its zeros. Biswas et al. also carried out numerical investigations on the 'amount of normalization and the extent of orthogonality' of the computed eigenfunctions in order to test their correctness. The overlap integrals required were evaluated by integration over x from  $-\infty$  to  $+\infty$ . We have seen in Section II.6 that any solution of the Schrödinger equation  $\psi(x;E) \rightarrow +\infty$  or  $-\infty$ as  $x \rightarrow +\infty$ . unless E is exactly equal to an eigenvalue. Even a truncated expansion (truncated at a certain high index) gives a hump in the nonclassical region in the computed eigenfunction, where the actual eigenfunction decays monotonically. The overlap integrals evaluated by integrating over infinite limits are therefore inaccurate and the test of the eigenfunctions used by Biswas et al. is inconclusive.

TABLE (II.1); Stabilization of the computed eigenvalues (System; The Quartic nharmonic Oscillator)

Iteration number	Numbers of $\{\Delta_m(\Xi)\}$ required	Stabilized correction, SE	initial + OE (m)
$n=0$ $\lambda=1$	Binitial = 1.5	c, = 2.0	
Н	17	-0.1121	1.387 516 248 824 27
2	L -	0.428x10 <sup>-2</sup>	1,352 344 510 018 44
23	22	C.7152x1C-5	1.392 551 641 726 52
7	3,4	-0.1562x10-9	1.392 351 641 530 29
ın	17	C.2538zl0-16	1.352 551 641 530 29
n=10 λ=10υC	C Enitiel=502,9864	c = 27.86	
7	55	-0,1601	502.836 327 005 776
2	44	0.7228x10 <sup>-4</sup>	502.386 359 286 167
27	50	C.1451x10-8	502.836 399 284 716
4	89	0.5718x10 <sup>-15</sup>	502.886 399 284 716
1			

TABIE (II.1)(...Contd.); Stabilization of the computed eigenvalues. (System: The Quartic Anhamonic Oscillator)

GAS.

Iteration number	Numbers of $\{\Delta_{m}(\Xi)\}$ required	Stebilized correction, Ja	Linitiel r off (m)
[0]0 0-1	E 2154.453	o• । ।	
1000 10-V 0001=II			
Н	1682	-0.2050	2 134.244 535 855 36
7	1700	-C.1586x10 <sup>-2</sup>	2 154.242 547 928 25
	1725	-0.2694x1c-5	2 154.242 545 254 23
4	1744	-0,2018x10 <sup>-8</sup>	2 154.242 545 252 21
ΓC	1764	-0.1512x10 <sup>-11</sup>	2 154.242 545 252 21
n=1600 >=40vic	Binitial = 747 785.382	a = 44.2.83	
7	1681	0.3871x1c <sup>-1</sup>	747 783.421 517 234
2	1657	-0.1440x10 <sup>-4</sup>	747 783.421 502 856
tr\	1713	-0.3001x10 <sup>-8</sup>	747 733.421 502 854
<i>\\</i> -	1718	0.9092x10 <sup>-10</sup>	747 787.421 562 834
			t-st. of the de to the foregree of the descriptions and the description of the the the total of the the total of the total

TABLE (II.2) : Comparison of our Results for E (  $\lambda = 1$ ) with the Results of Earlier Calculations.

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Quantum number	$E_n$ ( $\lambda = 1$ ) for the Quart	ic Anharmonic Oscillator
n	This work	Earlier works 1 392 35 164 1.387 187 78 (a)
0	1.392 351 641 530 29	1.392 351 641 550 29 (b) 1.392 350 653 679 1 (c)
10	53.449 102 139 665 3	53.448 404 6 (d)
100	1 035.544 183 138 91 .	1 035.544 04 (d)
1000	21 932.783 710 666 9	21 932.783 6 (d)
10000	471 103.777 790 809	471 103.778 (d)
	$E_n$ ( $\lambda = 1$ ) for the Pure	Quartic Oscillator
	This work	Earlier works
U	1.060 362 090 464 18	1.060 362 090 48 (e)
10	50.256 254 516 682 9	50.256 254 516 7 (e) 50.256 254 0 (d)
100	1 020.989 992 105 37	l 020.989 99 (d)
1000	21 865.262 118 137 7	21 865.262 1 (d)
10000	470 790.294 427 023	770 790.293 (d)

<sup>(</sup>a) Hioe and Montroll (1975) - Egn. (III.9).
(b) Biswas et al. (1973).
(c) Graffi et al. (1969).
(d) Hioe and Montroll (1975) - sophisticated WKB values.
(e) Reid (1970).

TABLE (II.3): Eigenvalues of the Pure Quartic Oscillator  $(H=p^2+\lambda x^4) \text{ and the Quartic Anharmonic Oscillator } (H=p^2+x^2+\lambda x^4) \text{ for } \lambda=1.$ 

Quantum number n	Pure Quartic Oscillator Eigenvalues	Quartic Anharmonic Oscil- lator Eigenvalues
O	1.060 362 090 484 18	1.392 351 641 530 29
1	3.799 673 029 801 40	4.643 812 704 212 08
2	7.455 697 937 986 74	8.655 049 957 759 31
3	11.644 745 511 378 2	15.156 803 898 049 9
4	16.261 826 018 850 2	19.057 557 436 303 3
5	21.238 372 918 236 0	23.297 441 451 225 2
<b>ં</b>	26.528 471 183 682 5	28.835 338 459 504 2
7	32.098 597 710 968 3	34.640 848 321 111 3
8	37.925 001 027 034 0	40.690 386 082 106 4
9	43.981 158 097 289 7	46.965 009 505 675 5
1.0	50.256 254 516 682 9	55.449 102 139 665 3
11	56.734 214 055 173 0	60.129 522 959 157 8
1.2	63.403 046 986 718 9	66.995 030 001 247 2
13	70.252 394 628 616 6	74.035 874 359 102 5
1.4	77.273 200 481 984.0	81.243 505 050 767 2
15	84.457 466 274 942 0	88.610 348 800 799 2
16	91.798 066 808 991 2	96.129 642 045 234 1
17	99.288 606 660 493 3	103.795 300 322 273
18	106.923 307 381 733	111.601 815 045 173

Table (II.3) (..Contd.)

Quantum number n	Pure Quartic Oscillator Eigenvalues	Quartic Anharmonic Uscil- lator Eigenvalues
1.9	114.696 917 384 985	119.544 170 733 050
20	122.604 639 000 999	127.617 777 795 355
21	130.642 068 748 630	135.818 417 325 610
22	138.805 147 911 395	144.142 195 296 398
23	147.090 121 257 604	152.585 504 205 574
24	155.493 502 268 682	1.61.144 990 694 51.3
25	164.012 043 622 865	169.817 528 001 595
26	172.642 711 962 845	178.600 192 366 876
27	181.382 666 185 768	187.490 242 692 950
28	190.229 238 652 463	196.485 102 910 221
29	199.179 918 833 747	205.582 346 604 423
20	208.232 339 005 144	214.779 683 549 177
31	217.384 261 674 103	224.074 947 852 600
32	226.633 568 481 138	233.466 087 479 375
<b>3</b> 5	235.978 250 361 696	242.951 154 951 147
34	245.416 398 791 936	252.528 299 061 493
35	254.946 197 970 798	262.195 757 468 520
36	264.565 917 814 499	271.951 850 050 007
37	274.273 907 658 941	281.794 972 923 820
38	284.068 590 581 401	291.723 593 051 013

TABLE (II.3) (...Contd.)

Quantum nu	mber Purc	Quartic ( Eigenva		· ·	Anharmor or Eiger	
39	293.9	48 458 20	66 006	301.736	243 351	187
40	305.9	12 066 3	48 384	311.831	518 269	701
41	313.9	58 030 1	83 978	322.008	069 744	845
42	324.0	85 020 99	92 133	332.26 <i>4</i>	603 530	091
43	334.2	91 762 3	34 482	342.599	875 832	5.47
44	344.5	77 026 8	91 585	357.012	690 233	780
45	354.9	39 633 5	06 395	363.501	394 863	479
46	365.3	78 444 4	67 063	374.066	J79 800	092
47	375.8	192 363 0	04 953	384.705	074 675	721
48	386.4	.80 330 9	86 517	395.416	946 465	263
49	397.1	41 326 7	80 67 <i>4</i>	.106.200	997 442	128
50	4.07.8	374 363 2	84 438	417.056	263 284	848
						ŧ

÷ MABLE (II.4) : The Quartic Anharmonic Oscillator Eigenvalues for Various values of

n		NAME, P. P. P. CHARLES OF THE THEOREM THE CHARLES OF THE THE THE CHARLES OF THE THEOREM CHARLES OF THE CHARLES	
10000.	1.000 007 499 358 76	3.000 037 498 968 31	5.000 057 496 155 56
.0001	1,000 074 986 88C 2C	5,000 374 895 936 12	5.000 374 615 958 59
.001	1.000 748 692 573 19	3.063 739 748 163 73	5.009 711 872 788 11
. 03.	. 1.067 375 672 081 58	3.036 525 304 513 35	5.095 959 132 742 51
-	1.655 285 509 543 72	3,506 872 013 152 92	5.7:7 959 268 833 56
1.0	1.392 351 641 550 29	4.548 812 704 212 08	3.555 049 957 759 31
10	2,449 174 072 118 39	8,555 003 454 807 77	15.655 521 492 413 8
100	4.995 417 545 137 59	17.830 192 715 952 5	54.675 384 261 994 8
1000	10,659 788 711 526 1	38.086 833 459 382 3	74,581 404 200 164 8
10000	22,861 608 870 272 5	81.503 316 953 284 5	160,565 912 611 712
40000	36.274 458 155 736 8	129,973 351 403 294	255.017 677 285 574

TABLE (II.4) (... Contd.) : The Quartic Anharmonic Oscillator Figurelues for Various Values of A.

u	3	4	3
A The same of the same of	Willer impose o excelemental emission of a view with the enterior of the second of the	A ANY PROCESSIONAL PROCESSION OF THE PROCESSION OF THE PROCESSION OF THE RECEIPED THE TAX CHIEFLY OF THE PROCESSION OF T	
.00001	7.00c 187 496 157 29	9.066 307 479 696 43	11.0vv 457 465 499 5
.0001	7.001 874 016 667 66	9.003 072 572 244. 61	11.064 571 355 129 7
.001	7.018 652 592 657 52	9.030 549 536 074 71	11.045 590 587 179 5
.01	7.178 573 180 700 50	9.289 479 815 311 89	11.425 792 646 186 3
Ļ	8.552 677 825 755 75	11.098 595 522 633 0	13.969 926 197 742 8
1.0	13,156 803 893 049 9	18.057 557 436 303 3	23.297 441 451 223 2
10	25.806 276 215 055 7	55.385 171 222 253 9	46.725 680 966 817 1
100	54.385 291 571 605 1	75,877 004 628 659 7	95.032 857 315 407 5
1000	116.603 198 937 293	162.802 374 195 975	212.554 185 409 754
,10000	250.950 743 891 715	350.435 896 215 566	457.654 575 005 690
4.0000	398,250 246 556 059	555.200 474 650 524	726.405 686 448 353

AABLE (II.4)(...Contd.) : The Quertic Anharmonic Cacillator Eigervalues for Various

T		T. 1	8
7	Anderson the state of the state		
רטטטט	15,000 657 440 292 3	15.000 347 403 300 5	17,001 687 567 750 3
1000	15,636,569,351,227	15.008 465 397 365 C	17.010 861 805 528 7
T000.	153	15.085 856 587 626 Û	17.107 457 792 553 5
10.	715	15,771 515 065 642 6	17.979 510 583 711 2
	16.954 794 686 144 1	20,043 863 604 188 5	23.229 552 179 939 3
1.0	28,835 338 459 504 2	54.540 848 321 111 3	40.590 586 082 106 4
0.0	58,241 298 759 755 2	70,351 051 939 234 7	83.005 867 037 585 3
100	123.540 657 625 578	149.545 557 443 288	175,628 655 957 714
1000	265,519 951 678 280	321,244 760 274 355	379.511 311 178 729
10000	571,547 791 619 426	691,663 457 635 372	817.156 874 968 737
70007	907.329 749 584 590	1 097.832 281 315 18	1 297.050 657 027 22

TABLE (II.4)(...Corti.) : The Quartic Arharmonic Cscillawor Ligenvelues for Various .

d/	5			10		
-	entrantiamente estructural de l'action de l'action de la company de la company de la company de la company de l					
.00001	19,601 557 515 867	7	21,667	. 557 251	1 378	22
.0001	19,013 556 530 636	7	21,016	550 255	5 642	2
.001	19.133 955 491 852	71	21.163	358 105	5 765	w
.61	20,210 670 452 755	ں	22,462	: 505 5-2	2 166	2
	26.505 554 752 536	9	29, 866	525 234	4 671	10
0.1	46.965 009 505 675	ピノ	55.449	102 139	9 655	1.7
10	56.156 252 981 197	7	109,772	570 864	333	
700	204,794 774 512 945		255,956	225 876	5 276	
700C	440.114 532 253 656		502.885	399 284	972 1	
10000	947.685 951 666 079		1 082,883	518 002	37 3	
40000	1 504.225 045 052 05		1 718,854	435 887	80 /	

TABLE (II.4)(...Contd.) : The Quartic Anharmonic Oscillator Eigenvalues for Various values of  $\lambda$  .

п			10.5				7,000	0C		
.00001		201.151	252	371	240	2	015.807	853	4.15	28
.0001		202,494	610	593	242	2	134.242	545	252	21
.001		214,458	455	291	852	2	316.405	517	618	20
TO.		285.365	C7C	205	075	5	019.741	205	332	61
** 		504.855	925	630	L50	10	294,061	322	693	S
1.0	H	055.544	185	138	16	27	932.785	710	999	٥٦
10	N	206,428	699	064	32	47	138.656	955	225	$\sim$
100	4	77.2.160	515	320	88	101	504.115	156	550	
76.00	10	211.359	950	955	0	218	655.377	196	231	
10000	23	997,240	274	526	G)	471	075.928	575	257	
40000	34		4.93	777	۲	727	785.;21	502	834	
								İ		

TABLE (II.5): Eigenvalues of the Quartic Anharmonic Oscillator  $(\pi = \rho^2 + x^2 + \lambda x^4) \text{ in Regimes of Extreme Values of } (n, \lambda).$ 

n	0.0001	40000 
O	1.000 074 986 880 20 (near harmonic regime)	36.274 458 133 736 8 (near quartic regime)
1.000	2 134.242 545 232 21 (boundary layer)	747 785.421 502 834 (near quartic/WKB regime)

 $\zeta$ .

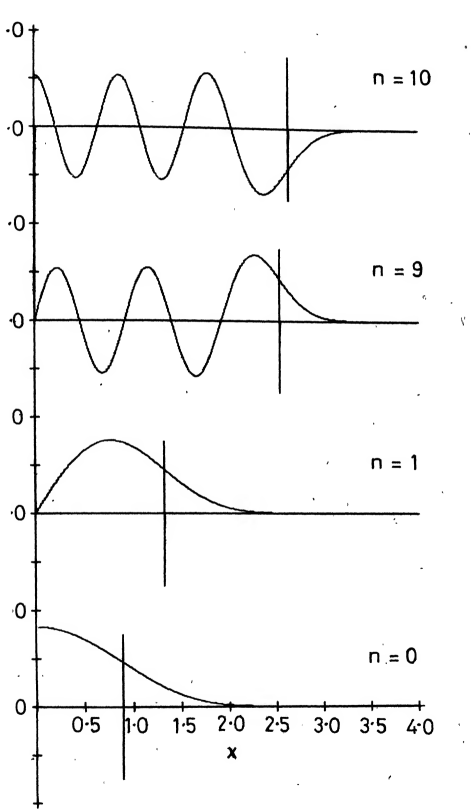


Fig. II.1 Quartic Anharmonic Oscillator Eigenfunctions for  $\lambda = 1$ 

#### CHAPTER III

#### THE CENERAL ANHARMONIC OSCILLATOR

### III 1 Introduction

The cigenvalue problem of the general anharmonic oscillator described by the Hamiltonian

$$H = p^2 + x^2 + \lambda x^{2\mu}, \qquad (5.1)$$

where  $\lambda > 0$ ,  $\mu = 3,4$ , ... and  $p = -i \frac{d}{dx}$ , is considered in this chapter. Using the realing arguments similar to those in Chapter I, it follows that the eigenvalues of  $H(k,\lambda) = \mu^2 + kx^2 + \lambda x^{2\mu}$  are given by  $E_n(k,\lambda) = k^2 L_n(1,\lambda^2)$ , where  $\lambda^2 = k^{-3/2}\lambda$ . It ensures that the eigenvalue problem of the anharmonic oscillator  $H(k,\lambda)$  can be completely described in terms of the reduced Hamiltonian  $H(1,\lambda)$ .

A straightforward perturbative solution of this problem runs into difficulties. The perturbation expansion for the eigenvalues in powers of  $\lambda$  is not convergent but asymptotic (bimon 1970). The coefficients in the perturbation series grow very fast and the construction of various Padé approximants become extremely involved. The numerical results of Craffi et al. (1971) for the octic anharmonic oscillator

(µ=4 in (3.1)) suggest that Pade approximents do not converge to the exact eigenvalue. However, mixed Borel-Pade method were utilized to obtain a few eigenvalues. Non-perturbative calculations have been relatively more successful. et al. (1973) extended the 'Hill determinant' method, used for the quartic anharmonic oscillator problem, for this case. whey used the basis functions  $\{x^m e^{-x^2/2}\}$  for the expansion of the eigenvalues for the sextic and the octic anharmonic oscillators ( $\mu$ =3 and 4 respectively in (5.1)) for values of  $\lambda$  in the range  $0 < \lambda \le 100$  for n=0 and  $0 < \lambda \le 10$  for n=2. However, the accuracy of the computed eigenvalues in their vork reduces significantly as one goes from the quartic to the sextic or the octic anharmonic oscillator, besides, the evaluation of the eigenvalues gets confined to smaller regime of  $(n,\lambda)$  values. Lakshmanan and Prabhakaran (1973) obtained comiclassically an asymptotic expression for  $\mathbf{E}_{n}^{(2\mu)}(\boldsymbol{\lambda})$  in the μ=3 case. Truong (1975) used Weyl-quantization prescription to study the sextic anharmonic oscillator eigenvilues but no new regults were obtained. For sufficiently large n, the WKB approximation method has been used to obtain approximate eigenvalues.

The most recent and extensive work on the general anharmonic oscillator eigenvalue problem is due to Hioe, MacMillen and Montroll (1976). They distinguished two limiting regimes

of values of  $(n,\lambda)$  for the eigenvalues  $\mathbb{E}_n^{(2\mu)}(\lambda)$  analogous to the similar distinction made for the quartic anharmonic oscillator eigenvalues. In one regime the energy eigenvalues differ slightly from the harmonic oscillator levels (the hear harmonic' regime); in the other they differ slightly from the pure 2p-ic oscillator eigenvalues (the 'near pure anharmonic' regime). The above two limiting regimes are separated by a regime called the 'boundary layer' in which the energy eigenvalues are not 'nearly harmonic' or 'nearly pure anharmonic'. Hioe et al. used different formulations of the cigenvalue problem in various regimes and constructed deveral simple formulae with different ranges of validity. Using Bargmann representation, they developed numerical algorithms from which the energy eigenvalues in the small n regime may be computed. The algorithm is similar to that developed for the quartic anharmonic oscillator eigenvalue problem. First few eigenvalue: were thus computed to 5-6 significant figures for the sextic and the octic anharmonic oscillators, for various value of  $\lambda$ . However, the size of the determinants, required for the computation of the eigenvalues, increases rapidly with  $\mu$  or n, making the evaluation of higher eigenvalues laborious.

We show in this chapter that the method described in Chapter II may be extended to solve the general anharmonic

oscillator eigenvalue problem and eigenvalues of arbitrarily high accuracy are obtainable in this case in all regimes of  $(n,\lambda)$  as for the quartic anharmonic oscillator problem.

# III.2 The Appropriate Scaling Formula

The eigenvalues  $E_n^{(2\mu)}(\lambda)$  and the eigenfunctions  $\psi_n^{(2\mu)}(x;\lambda)$  of a general ahnarmonic oscillator (H = p<sup>2</sup> +  $x^2 + \lambda x^{2\mu}$ ) are the solutions of the Schrödinger equation:

$$\left[-\frac{d^{2}}{dx^{2}}+x^{2}+\lambda x^{2\mu}\right]\psi_{n}^{(2\mu)}(x;\lambda) = \mathbb{E}_{n}^{(2\mu)}(\lambda)\psi_{n}^{(2\mu)}(x;\lambda),$$
(3.2)

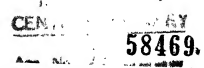
with the boundary condition  $\psi_n^{(2\mu)}(x;\lambda) \to 0$  as  $x \to \pm \infty$ . We write eigenfunctions in the form

$$\psi_{n}^{(2\mu)}(x;\lambda) = e^{-\alpha x^{2}} \sum_{m=0}^{\infty} a_{m} x^{m},$$
(3.3)

where  $\alpha$  is the scaling constant. A formula for determining the appropriate scaling  $\alpha$  for any  $\mu$  is obtained along the same lines as for the quartic anharmonic oscillator (section II.1). The exact n-th eigenfunction has n zeros in the region of oscillation which, for sufficiently large n, is  $\sqrt{(E_n^{-(2\mu)}/\lambda)^{1/2\mu}}$ . In the WKB approximation

$$\mathbb{E}_{n}^{(2\mu)}(\lambda) \simeq C \lambda^{\frac{1}{\mu+1}} \left(n + \frac{1}{2}\right)^{\frac{2\mu}{\mu+1}}.$$

Hence, the region of oscillation  $\sqrt{\lambda} = \frac{1}{2(\mu+1)} \frac{1}{(n+\frac{1}{2})^{(\mu+1)}}$ . For an effective expansion the region of oscillation must



include the span of first n (or a number proportional to n) basis functions. The appropriate scaling formula is thus obtained by setting the region of oscillation of the nth eigenfunction equal to the width of the nth basis function ( $^{\circ}\alpha^{-1/2}$   $n^{1/2}$ ), which yields

$$\sigma(n,\lambda) \sim (n+\frac{1}{2})^{\frac{\mu-1}{\mu+1}} \lambda^{\frac{1}{\mu+1}}$$
.

The above scaling formula is not expected to be good when n and/or  $\lambda$  in small, in view of the WKB approximation for  $E_n^{(2\mu)}(\lambda)$  used in the derivation. However, for small n, small  $\lambda$  the scaling must approach 1/2 — the appropriate scaling for the harmonic oscillator. Hence, the scaling formula valid in all regimes of  $(n,\lambda)$  and for any  $\mu$  is

$$\alpha(n, \lambda) = \frac{1}{2} + (n + \frac{1}{2})^{\frac{\mu-1}{\mu+1}} \lambda^{\frac{1}{\mu+1}}$$
 (5.4)

The rollowing regimes may be distinguished according to the above scaling formula:

$$(n + \frac{1}{2})^{\frac{\mu-1}{\mu+1}} \lambda^{\frac{1}{\mu+1}} \ll \frac{1}{2}$$
 is the near harmonic regime,

$$(n + \frac{1}{2})^{\frac{\mu-1}{\mu+1}} \lambda^{\frac{1}{\mu+1}} >> \frac{1}{2}$$
 is the pure anharmonic regime,

$$(n + \frac{1}{2})^{\mu+1} \frac{1}{\lambda^{\mu+1}} = \frac{1}{2}$$
 is the boundary layer between the above two regimes.

The value of the combination  $(n+\frac{1}{2})^{\mu-1}\lambda$  determines the regime to which the eigenvalue  $E_n^{(2\mu)}(\lambda)$  belongs. It may

be seen from above that the range of  $(n,\lambda)$  values for the 'near harmonic regime' dimnishes on increasing  $\mu$ . Hence the eigenvalues of general anharmonic oscillators, obtained from the methods which use a finite term expansion for the eigenfunctions in the harmonic oscillator basis  $\{x^m e^{-x^2/2}\}$ , deteriorate in accuracy on going to oscillators of higher  $\mu$ . The use of basis, appropriately scaled according to the regime, leads to a uniform treatment of the anharmonic oscillators eigenvalue problem for all u and in all regime of  $(n,\lambda)$ .

### III.3 Method

The expansion (3.3) on substitution into the Schrödinger equation for the general anharmonic oscillator (3.2) yields the following ( $\mu$ +2)-term linear recurrence relation among the expansion coefficients  $\left\{a_{m}\right\}$ :

$$(m+1)(m+2) a_{m+2} + (E-4\alpha m-2\alpha) a_m + (4\alpha^2-1)a_{m-2} - \lambda a_{m-2\mu} = 0$$
 (3.5)

The even and odd parity solutions are obtained respectively by assigning the initial conditions (i)  $a_0=1$ ,  $a_1=0$  (ii)  $a_0=0$ ,  $a_1=1$ . We divide the recursion (3.5) by (m+1)(m+2) and rewrite it in the following notations

$$a_{m+2} + d_{m,m} a_m + d_{m,m-2} a_{m-2} + d_{m,m-2\mu} a_{m-2\mu} = 0$$
,

where

$$d_{m,m} = \frac{(E-4\alpha m-2\alpha)}{(m+1)(m+2)}, \quad d_{m,m-2} = \frac{(4\alpha^2-1)}{(m+1)(m+2)}, \quad (3.6)$$

$$d_{m,m-2\mu} = -\frac{\lambda}{(m+1)(m+2)}.$$

For self consistency the determinant

The eigenvalues  $\mathbb{E}_n^{\ (2\mu)}(\lambda)$  are the roots of this transcendental equation. To obtain the roots numerically, we denote the determinant formed by omitting all rows and columns beyond  $d_{m,m}$  in  $\Delta(E)$  as  $\Lambda_{m+2}(E)$ . The determinant  $\Lambda_{m+2}(E)$  may be expanded into determinants of lower orders. It yields the following  $(\mu+2)$ -term recurrence relation among  $\{\Lambda_m(E)\}$ :

$$\Delta_{m+2}(E) - d_{m,m}(E) \Delta_{m}(E) + d_{m,m-2} \Delta_{m-2}(E)$$

$$+(-1)^{\mu-1} d_{m,m-2\mu} \Delta_{m-2\mu}(E) = 0.$$
(3.8)

Hence the values of the determinants  $\boldsymbol{\Delta}_{m+2}(E)$  upto any order

may be determined successively in terms of  $\Delta_0$  (or  $\Delta_1$ ) with the help of the above recursion. The corresponding zeros of  $\Delta_m(E)$ ,  $\Delta_{m+2}(E)$ ,... stabilize to an eigenvalue for large m provided the scaling  $\alpha$  is appropriate. The procedure for the actual numerical evaluation of the eigenvalues is the same as in described in section (II.2). The Newton's method, which is used for obtaining the roots of  $\Delta_m(E){=}0$ , requires the value of the derivative  $\Delta_m^*(E)$  also. The derivatives  $\Delta_m^*(E)$  can also be evaluated recursively with the help of the recursion:

$$\Delta_{m+2}^{i}(E) - d_{m,m}^{i} \Delta_{m}(E) - d_{m,m}(E) \Delta_{m}^{i}(E) + d_{m,m-2} \Delta_{m-2}^{i}(E) + (-1)^{\mu-1} d_{m,m-2\mu} \Delta_{m-2\mu}^{i}(E) = 0,$$
 (3.9)

obt ined by differentiating (3.8) with respect to E. The recursions (3.8) and (5.9) are continued on computer until the corrections for the required root of  $\{\Delta_m(E)\}$  for sufficiently large m stabilize to a prescribed extent. The eigenvalues accurate to 15 significant figures may thus be computed in only 4 or 5 iterations starting from rather crude initial estimates. The initial estimates for eigenvalues for large n (and not too low  $\lambda$ ) are obtained from the corresponding WKB approximation formulae and for low n they are obtained by evaluating a sufficiently large order determinant  $\Delta_m(E)$  from recursion (3.8) at various E points.

For obtaining eigenfunctions, E is set equal to the computed eigenvalues in the recursion (0.5) and the expansion coefficients  $\{a_m(E)\}$  are evaluated successively. The resulting coefficients  $\{a_m(E)\}$  provide a convenient representation for the corresponding eigenfunction through the expansion (0.5). The asymptotic behaviour of the solution of the Schrödinger equation (0.2)  $\exp(\pm |x|^{\mu^{+}/\mu + 1})$ , which requires

$$a_{m+2}/a_{m-2}$$
,  $\lambda/m^2$ ,  $m \to \infty$ , (5.10)

for the coefficients in the series solution (3.3). It is seen during the computation of  $\{a_m(E)\}$  that the above asymptotic dependence is satisfied actually, which ensures a decreasing (in magnitude) set of coefficients for sufficiently large m. It is now possible to obtain accurate eigenvalues and eigenfunctions for the general anharmonic escillators for any value  $(n, \lambda)$ .

# III.4 Eigenvalues

We have obtained the eigenvalues of the sextic  $(H = p^2 + x^2 + \lambda x^6, \lambda > 0) \text{ and the octic } (H = p^2 + x^2 + \lambda x^8, \lambda > 0) \text{ anharmonic oscillators for various values of n and } \lambda. The eigenvalues are computed using values of <math>\alpha$  in the range:

$$\alpha(n, \lambda) = \frac{1}{2} + (1.5 \text{ to 2.0}) (n + \frac{1}{2})^{1/2} \lambda^{1/4}$$
, (for the sertic anharmonic oscillator), (3.10a)

$$\alpha(n,\lambda) = \frac{1}{2} + (2.5 \text{ to } 3.0)(n + \frac{1}{2})^{3/5} \lambda^{1/5}, \text{ (for the octic anharmonic oscillator).}$$

The range of  $\alpha$  given above works for any value of  $(n, \lambda)$ . The eigenvolues, thus computed, are presented in tables (III.1) and (III.2) for n = 0, 1, 2, ... 10, 100 and 1000 and for different values of  $\lambda$  in the range .00001 $\leq \lambda \leq$  40000. they are evaluated to 16 figures and then are rounded off to 15 figures for the Mables. The Tables (III.1) and (III.2) include values of  $E_n^{(2\mu)}(\lambda)$  for the (high n, low  $\lambda$ ) region on the boundary layer. The computation of the eigenvalues in this region is found to be the most difficult in the ) on lier literature. Further, the values of  ${\mathbb E}_n^{\;\;(2\mu)}(\pmb{\lambda})$  for the sextic and the octic anharmonic oscillators to this accuracy in all regimes of (  $1,\lambda$ ) are regorted here for the first time the existing most extensive tables for  $E_n^{(2\mu)}(\lambda)$ , u = 3, 4, are due to Hioc et al. (1976) who have presented the values of first six eigenvalues to 6 significant figures for the sextic anharmonic oscillator and of first four eigenvalues to 5 significant figures for the octic anharmonic oscillator. The possibility of treating all higher order general anharmonic oscillators eigenvalue problem by the same technique for any value of  $(n,\lambda)$  is thus established in this chapter.

85 85 50 54 51 II  $\omega$ 0 \_ 718 634 301 189 553 351 008 655 519 726 251 305 784 TABLE (III.1) : The Servic Anharmonic Oscillator Figenvelues for Various Values SI7 015 703 656 108 301 911 525 711  $\sim$ 742 391 295 51,182 480 5,000 4,58 5.347 420 218 278 664 551 521 9,856 5.004 28.977 128.575 5,044 5,644 16.641 90.821 4.6 54 58 27 15 δĈ 70 2 4  $\sim$ ∵;-220 559 720 ST3 615 680 059 744 895 5,000 131 213 195 345 629 604 11.8 45.457 784 742 937 095 912 521 622 930 r 1 843 61,407 828 316 780 565 206 615 036 5,001 308 5,033 8,114 15.946 5.596 24.525 5.012 3,107 63. Ω Ω 35. 29 75 58 9 5  $\sim$ 7. LΩ 329 254 749 565 208 254 465 005 754 [5] [5] 572 27C 569 725 042 363 078 518 132 213 747 322 723 16.211 718 57.0 350 758 348 624 1,616 741 087 1,000 01.8 187 ں 3,716 11.478 2,205 6.492 1,169 1,435 7,000 1.001 40000 .00001 10000 .0001 1000 . CCI 100 1.0 . C7 70 -

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TABLE (III.1)(...Contd.) ; The Sextic Anhermonic Oscillator Eigenvalues for Varicus Values of  $\lambda$  .

u		4. }-	5
\ \	AND THE RESIDENCE OF THE PARTY		
.00001	7,601 100 315 756 71	5,102 415 887 905 VB	11.004 325 996 047 8
. 0001	7,011 720 523 720 43	9.025 St7 201 468 50	11.642 5u8 787 854 1
.001	7.116 692 855 856 92	9.218 581 748 732 25	11,377 CC8 617 207 5
.01	7.777 657 466 557 50	10,408 337 508 693 3	15.275 278 161 874 1
	10.237 873 721 423 9	14.507 64.0 046 120 3	1.2,861 758 353 358 2
1,0	15.989 440 787 825 7	22,916 186 450 728 6	56,622 590 570 533 5
70	27.155 085 614 651 4	35,289 550 657 370 3	52.849 512 578 258 2
700	47.564 984 581 593 5	69.046 575 526 034 7	93.073 891 695 377 6
1000	84.175 583 775 589 6	122,321 705 320 204	165.000 436 556 548
10000	145,457 970 316 556	217.261 599 779 375	253,120 422 079 698
40000	211,290 344 511 508	367.155 772 116 721	414.453 587 749 840

MABLE (III.1) (... Const.) : The Sertic unharmonic Oscillator Ligervilues for Various

n	9	7	ω
V	andriengalman manala. Itali dandak ang alika mininda angka manaka angka angka angka angka angka angka manaka m		
.0000	13.007 052 770 769 2	15.010 749 555 371 3	17.315 560 678 996 6
. 0001	13.059 151 375 425 9	15.104 797 508 119 9	17,150 733 586 946 5
.001	15,595 155 248 671 1	15.868 288 655 234 6	13,205 461 089 989 1
.01	16,25L 553 v42 653 7	13,445 522 881 748 1	22,312 380 711 280 6
H.	23.585 275 596 988 3	28,523 937 081 664 6	34.569 574 585 792 2
J, Č	39.051 906 854 586 1	43.141 505 571 449 5	57.845 728 456 525 1
10	67.632 071 647 819 2	83,730 950 282 579 2	100.865 986 116 256
100	119.399 778 956 875	147.857 547 351 650	178.239 969 897 024
1000	211.776 855 105 435	262,300 055 623 561	316,325 599 139 895
10000	576.275 559 274 236	455.107 291 369 825	552,158 183 022 017
40000	532.031 545 974 230	659.065 528 349 994	794.894 952 499 649

MABIR (III.1) (... Contt.) : The Sertic inharmonic Cscillator BiLenvalues for Valious

	9	ω	$\omega$	77	4.	1					
	213	557	24,1	957	435	484	455	563	546	455	14
10	252	616	451	218	318	575	922	577	101	18g	660
	loč	847	727	ú2ŭ	420	390	515	567	258	796	114
	21,029	21,277	25.070	36.032	46,508	78.558	135			455	275
	21,	21,	25.	30.	7.6	78.	158,135	244.478	434.045	771.455	096.875
											٦
	Н	-	<u>ي</u>	0	5	7					
	128	196	524	δόC	660	137	034	389	151	392	506
	325	5.53	378	025	950	875	924	157	4.06	116	561
())	571	.r20	977	475	244	291	133	618	593	4:2C	406
	15.021	19,203	20,505	26.345	40,408	68,128	050	437	575,653	047	958.981
	c., rd	J.c.	20.	26.	4°	68,	115.036	210,437	575,	554.047	978
	Ľ									C	0
l d	. JOCOI	. UCCI	, CO7	10.	٦	7.0	70	160	700C	τοοοο	40000
// ~	۰	۰	٥	•	•	П	П	1-1	• 1	, ,	7

TABED (III.1)(...Contd.); the Sextic arbarronic Oscillator Lisenvelues for Verious

	•	10										
	67.	e 15	11	0	5 1	9 9	!		10	_		
	ردي. دري	638	ζ), Γ'',	215	757	015	8	084	585	907	62	
コート	50.00	253	240	618	137	757	682	808	044	169	172	
Ä	515	373	725	511	635		362	523	534	285		
	545.575	545.878	847.725	724,511	541 (	700,650	480.952	684,523	101.534	825.235	739.9	
	4	2	12	22	40 ,	71	127 4	226	403 ]	3 91/	013 739.945	
							П	.,	7		7	
	4C4	378	205	342	52	63	71	10			•••	
	176 4	10 00 01	2 775	34C 3	522 7	442 (	465 7		168 7	826 2	821 8	
								, 611				
(C)	757	TSO	378	894	788	556	921	213	564	\$28	851	
	427	523	466	571	415	793	435	.,85	503	74.9	426	
	215.427	265.523	457.466	739.571	255,415	287,793	051,435	212.485	834.503	821.749	274.426	
				~	Н	~	4	_	12	22	32	
	)]	1								C	. `	
g/	, 00001	.0001	.001	10.	<b>~</b>	J.C	0.[	100	1000	10000	40000	
/ ~	,	•	•	-	•	П		<u></u>	<i>i</i> – 1	r1	7	

TABLE (III.2) : The Octic Anharmoric Oscillator Ligenvalues for Various Values of A.

61	99 851 081 375 600 3	001 017 100 100	5.025 554 559 087 81	5.190 981 556 774 52	5.880 4.8 512 335 72	7.559 948 490 535 78	16.993 737 535 505 0	15.711 022 781 994 9	26,055 458 321 253 1	40,974 759 857 386 5	64.760 471 754 927 2	1 552 512 566 587 1
	10 400 645 588 715 F	101 100	5.005 725 955 351 21	3.C47 977 747 255 24	5.268 551 508 038 18	5.939 721 551 641 99	5.368 778 061 748 15	7.929 683 682 350 75	12,195 021 933 650 2	19.090 Sl <sub>4</sub> 267 022 6	30.106 900 557 858 1	35.67C 505 9+5 098 3
(ب	000 000	1. UJO 055 520 ZIJ II	1.06C 646 359 374 C7	1,005 857 514 124 73	1.035 496 778 855 76	1.168 970 453 245 99	1,491 619 895 862 21	2,114 544 621 942 13	3,188 654 546 492 27	4.949 487 440 052 74	7.778 272 214 511 10	10,238 868 255 479 1
51/~		TOOOJ.	.0001	.001	.01	H.	٦. (	10	100	1000	10000	40000

TABLE (III.2) (... Corte.) a the vetic inhermonic Oscillator Ligenvelues for Various Talues of  $\lambda_*$ 

g/	17	4	5
· ·			
.0000	7,008 365 664 175 81	5.426 553 496 754 59	11, 7 522 812 186 7
,000	7,075 568 572 662 77	5.1E0 256 740 lus Sl	11,555 154 415 293 5
.001	7.507 561 558 254 58	10,045 355 306 563 2	12.021 759 638 194 8
.01	8,955 834 568 107 55	12,534,709,515,209,4	16.655 618 992 657 8
۲.	12,281 167 732 276 1	17.761 215 567 788 7	23.597 (86 021 463 5
1.0	18,191 100 018 514 9	26.743 448 558 041 2	56.509 236 308 241 5
10	28.022 750 232 932 1	41,494 702 572 696 9	56.858 550 129 970 3
100	43.902 11.3 335 199 5	65.201 815 832 253 8	39.569 748 759 825 3
1000	69,257 537 833 105 0	102,982 586 803 735	141.574 025 285 519
J.CC00	109,562 182 579 437	162.992 157 936 368	224.137 086 582 704
40000	144.492 517 309 256	214.555 131 730 875	295.650 523 015 270

TABLE (III.2)(... Conto.) ; The vetic anharmonic Secillator Eigenvalues for Various

&		17,219 185 575 051 6	18.455 698 918 205 9	22,558 588 757 649 3	51.287 557 841 939 9	46.726 039 572 425 5	72,241 657 072 499 5	115,548 652 162 102	178,921 526 920 463	283,114 615 544 210	448,418 117 625 494	591.584 145 926 580
· ·		15.157 771 538 185 8	15.984 519 524 996 2	19,684 413 273 066 1	26.019 385 438 593 8	38,519 256 955 771 0	59.323 544 225 652 5	92.930 664 512 525 0	146.596 403 041 015	231,96.: 927 760 861	367.270 224 518 525	484,514 056 324 413
9		2 13 620 773 670 261 5	420 365	15.636 253 113 142 6	21,155 638 455 788 5	30,951 097 089 331 5	47.593 375 092 009 1	74,083 047 442 977 1	116.762 998 351 560	164.646 163 263 619	292,385 266 491 847	565,708 784 555 478
U/	/ /	Light Act of	10000.	100.	.01		1.0	10	100	1000	10000	46600

TABLE (III.2)(... Contd.) : The Cotic Arkarronic Cscillator Digenvalues for Various

	C7	1.0
. 00001	10,529 759 685 555 1	21,475 815 823 694 1
. c.cl	21,035 500 450 453 8	25,728 451 754 605 3
	25,251.586 225 766 0	30.155 815 137 584 6
	76,921 273 635 926 1	42.905 502 950 440 2
	55.522 679 758 570 0	64.882 127 949 355 0
	86.099 541 860 127 4	160.856 404 455 769
	155.259 874 530 135	158.599 173 255 695
	213,615 701 595 858	250.575 151 337 872
7007	538.686 489 508 001	396.637 899 756 OL5
10000	535.520 106 526 578	628.315 087 274 892
40000	117 521 160 115,307	828.950 218 446 123

TABE? (III, 2)(... Contd.); Te Cotic . Therronic Cacillator Digentalues for Various

1 477.162 84c 775 5c       58 c35.255 275 976         2 535.412 136 221 36       52 c39.504 286 6         5 555.528 705 597 06       145 660.545 560 5         5 850.452 007 751 74       231 165.653 875 4         9 270.398 457 27c 55       556 568 0;1 845 5         14 691.365 575 93c 8       580 651.053 558 5         23 285.471 395 490 2       920 266.633 525 5         30 722.437 520 580 8       1 214 500.225 845 36	.000Cl .CCCl	., 9	4C1.157 605.179 979.755	100 7 (25) 9 115 5 521	304 567 528	53 711. 166	25.2	10 656,172 107,125 568,700	1000 172 S 125 C 107 7	51.5	657 525 772	4 01 0)
2 535.412 136 221 38	, ,		.77.162			56	58		523	275	545	ω
3 595,528       703 597 06       145 660,545 560         5 850,452 007 751 74       231 165,653 875         9 270,598 457 270 55       556 568 071 845         14 691,365 575 930 8       580 651,093 558         23 285,471 395 490 2       920 263,635 625         30 722,437 520 580 8       1 214 500,225 845			35.41		21	38	35			286	500	
5 850.452 007 751 74       251 165.565 875         9 270.598 457 270 55       556 568 0.71 845         14 591.565 575 930 8       580 651.053 558         25 285.471 595 490 2       920 260.033 525         30 722.437 520 580 8       1 214 500.225 845	,		195.528			90	145			560	585	
9 270, 598 457 270 55 568 0.;1 845 14 691, 365 575 930 8 580 651, 093 558 23 285, 471 395 490 2 920 260, 033 525 30 722, 437 520 580 8 1 214 500, 229 845	Δ,		50.45			74	251	165,6		875	444	
14 691.365 575 930 8       580 651.053 558         23 285.471 595 490 2       920 268.635 525         30 722.437 520 580 8       1 214 500.225 845	01		70.39			35	556				252	
23 285.471 395 490 2 920 260.633 525 30 722.437 520 580 8 1 214 500.229 845	77		91.36			ω	580	651.0			558	
30 722,437 520 580 8 1 214 500,229 845	10000		85.47.		96	(A	920				588	
	4000C 30		22,43					500.2			36	

#### CHAPTER IV

### THE DOUBLE MINIMUM OSCILLATOR

## IV.1 Introduction

We consider in this chapter the eigenvalue problem of the double minimum oscillator (d.m.o.) described by the Hamiltonian

$$II(l_{\lambda}) = p^2 - x^2 + \lambda x^4, \quad \lambda > 0. \tag{4.1}$$

The potential function of a d.m.o. has two symmetric potential wells separated by a barrier. A feature of its eigenvalue problem is the bunching of the lower eigenvalues in pairs for sufficiently large separation between the two wells. The d.m.o. models some interesting physical problems. The vibilitional spectra of some releculer possess two parallel type nearly superimposed bands, a phenomenon which may be directly related to the eigenvalue spectrum of the d.m.o. The commonly known example in this regard is the inversion spectra of the ammonia molecule (see Domnison and Uhlenbeck 1952). Besides, the potential functions of several hydrogen bonded solids are found to possess two minima in the region available for protonic movement (see Synder and Ibers 1962,

Somorjai and Hornig 1962). The most recent appearance of this model is in the spontaneous symmetry breaking (Polyakov 1977).

The computation of the splitting between the eigenvalues forming pairs has been the subject of considerable interest. The aplitting depends in general on the separation between the two wells and the nature of barrier between them. d.m.o. with potential function  $V(x) = \frac{1}{2}k (|x| - a)^2$  is exactly solvable (Merzbacher 1961) and an expression for the splitting between the two lowest eigenvalues is  $\Delta E \sim k^{2}a \exp(-ka^{2})$ . Dennison and Uhlenbeck (1932) obtained the splitting in the KB approximation and then compared the UKB values with the exact values for a d.m.o with potential function formed by joining two equal parabolae with a straight ling. The WKB values for the splitting are found fairly accurate for large separations between the two parabolae. It is interesting to note that the WKB approximation is applied here for low n. the usefulness of those results is, however, limited to some extent, as the potential functions used in these works are non-analytic. Harmony (1971) trusted the dim.o. problem vii a harmonic oscillator approximation and obtained zeroorder and first-order expressions for the splitting.

The perturbation expansion of the energy eigenvalues of the d m.o. described by (4.1) in power series of  $^\lambda$  is

Non-convergent (Simon 1970). Somorphi and Hornig (1962) obtained numerically a few energy eigenvalues (to 5-5 significant figures) for the d.m.o. with Hamiltonian  $H(k,\lambda) = p^2 + kx^2 + \lambda x^4$  for five different pairs of values of  $(k,\lambda)$ . The calculations were denotely expanding the eigenfunctions in the harmonic oscillator basis functions and diagonalizing the secular determinant formed. We obtain, in this chapter, accurate eigenvalues and eigenfunctions of (4.1) using the method described in Chapter II. A VLB expression for the splitting is also obtained for this problem and the VKB values are compared with the corresponding accurate values for the splitting for various values of  $\lambda$ .

## IV.2 Eigenvaluos

The ochrödinger equation for the d.m.o. (4.1) is

$$\left[-\frac{d^2}{dx^2} - x^2 + \lambda x^4\right] \psi_n(x;\lambda) = E_n(\lambda) \psi_n(x;\lambda) . \quad (4.2)$$

The digenfunctions are expanded as

$$\psi(\mathbf{x}; \boldsymbol{\lambda}) = e^{-\alpha \mathbf{x}} \sum_{m=0}^{\infty} a_m \mathbf{x}^m, \qquad (4.3)$$

which on substitution into (4.2) yields the following 4-term recurrence relation,

 $a_{m+2} + d_{m,m}a_{m} + d_{m,m-2}a_{m-2} + d_{m,m-4}a_{m-4} = 0,$  (4.4) Where

$$d_{m,m} = \frac{(E - 4\alpha m - 2\alpha)}{(m+1)(m+2)}, \quad d_{m,m-2} = \frac{4\alpha^2 + 1}{(m+1)(m+2)},$$

$$d_{m,m+4} = -\frac{\lambda}{(m+1)(m+2)}.$$

The method of computing the eigenvalues from a recursion of the type (4.4) has been described in Chapter II. Therefore, the eigenvalues of the d.m.o. are obtainable with uniform accuracy in all regimes of  $(n,\lambda)$ . The characteristic bunching of the eigenvalues in pairs occurs for small n and for sufficiently large separation between the two wells. Since the separation between the two wells  $\sqrt{(1/\lambda)}$ , the region of interest for the present problem is the (low n, low  $\lambda$ ) regime. For (high n, high  $\lambda$ ) regime the eigenvalues are near pure quartic. We have therefore computed the cigenvalues in the (low n, low  $\lambda$  ) regime and the results are presented in Table (IV.1) for the first eight eigenvalues for values of  $\lambda$  in the range (.01  $\leq \lambda \leq$  0.20. values of the scaling α used in these computations lie between 0.5 and 1.0. The eigenvalues presented in the Mable (IV.1) are with respect to the bottom of the potential wells at zero energy and is related to  $\mathbf{E}_{\mathbf{n}}(\lambda)$  by

$$\varepsilon_{n}(\lambda) = \frac{1}{4\lambda} + E_{n}(\lambda),$$
 (4.5)

where  $1/4\lambda$  is the depth of the potential well. The numbers  $\varepsilon_n(\lambda)$  are positive definite and provide a direct look at the variation of the eigenvalues with  $\lambda$ . For  $\lambda \neq 0$ , the deparation between the two wells increases and the probability of penetration through the barrier approaches zero. The eigenvalues  $\varepsilon_{2n}$  and  $\varepsilon_{2n+1}$ , therefore, become nearly doubly degenerate for small n. For instance,  $\varepsilon_{2n}$  and  $\varepsilon_{2n+1}$  (n = 0,1,2) are found close to each other to at least 14 significant figures for  $\lambda$  =0.01. The expansion of the potential function of the d.m.o. about the minima of the well  $\sqrt{2}x^2$ , for  $\lambda \neq 0$ ; therefore the lower eigenvalues  $\sqrt{2}(2n+1)$ . The numerical results confirm this observation.

The eigenvalues of  $H(k,\lambda)=p^2-kx^2+\lambda x^4$  are obtainable from the eigenvalues of  $H(1,\lambda')$  using the scaling relation

$$\mathbb{E}_n(\mathbf{k},\lambda) = \mathbf{k}^{1/2} \, \mathbb{E}_n(\mathbf{l},\lambda'),$$
 where  $\lambda' = \mathbf{k}^{-3/2}\lambda$ .

# IV. 3 The WKB Formula for Splitting

The splitting between the pairs of lower eigenvalues of a symmetric d.m.o. in the UKB approximation is given by (Landau and Lifshitz 1965)

$$\Delta E^{VKB} = \frac{\omega}{\pi} \exp \left[ - \int_{-x_{O}}^{x_{O}} |p| dx \right], \qquad (4.6)$$

where  $\omega^{-1} = \frac{1}{2\pi} \int_{x_0}^{x_1} p^{-1} dx$  and  $\pm x_0$ ,  $\pm x_1$  are the four turning

points. The derivation of the above formula assumes small probability of penetration through the barrier. For the d.m.o. described by (4.1) the turning points are given by

$$x_0^2 = \frac{1}{2\lambda} (1-u), \quad x_1^2 = \frac{1}{2\lambda} (1+u),$$
 (4.7)

where  $u = \sqrt{(4 \lambda \epsilon_n^0)}$ ,  $\epsilon_n^0 = \frac{1}{4 \lambda} + E_n^0$ ,  $E_n^0$  is the mean energy of the two eigenvalues forming a pair. The integrals involved in (4.6) may be expressed in terms of the complete elliptic integrals K(k) and E(k) of the first and the second kinds respectively (Gradshteyn and Ryzhik 1965)

$$\int_{-x_0}^{+x_0} |p| dx = \lambda^{1/2} \int_{-x_0}^{+x_0} [(x_0^2 - x^2)(x_1^2 - x^2)]^{1/2} dx$$

$$= 2\lambda^{1/2} \frac{x_1}{3} [(x_0^2 + x_1^2) E(t) - (x_1^2 - x_0^2) K(t)] (4.8a)$$

and

$$\int_{x_{0}}^{x_{1}} p^{-1} dx = \frac{1}{\lambda^{1/2}} \int_{x_{0}}^{x_{1}} [(x^{2}-x_{0}^{2})(x_{1}^{2}-x^{2})]^{-1/2} dx$$

$$= \frac{1}{\lambda^{1/2}x_{1}} K(q) , \qquad (4.8b)$$

where  $t = (\frac{1-u}{1+u})^{1/2}$ ,  $q = (\frac{2u}{1+u})^{1/2}$ . Thus,

$$\Delta E^{\text{VKB}} = \frac{2^{1/2}(1+u)^{1/2}}{I(q)} \exp \left[-\frac{2^{1/2}}{5\lambda} (1+u)^{1/2} \left(E(t) - uL(t)\right)\right]$$
(4.9)

The values of the splitting AE are calculated from the above formula for various values of  $\lambda$  for the lowest two eigenvalues

and are compared with the corresponding accurate values in Table (IV.2). The mean energy  $\mathbb{D}_n^0$  required in (4.9) is evaluated from Table (IV.1). The WKE values for splitting are surprisingly good.

For small  $\lambda$  a simple analytic approximate expression for AE may now be obtained from (4.9) using the following expressions for the elliptic integrals (Gradshteyn and Ryzhik 1965):

$$E(k) = \frac{\pi}{2} (1 + \frac{1}{4} k^2) + O(k^4),$$

$$E(k) = \frac{\pi}{2} (1 - \frac{1}{4} k^2) + O(k^4), \quad k \to 0$$
(4.10)

and - 4 1 - 4 - 2 - 2

$$K(k) = \ln \frac{4}{k!} + \frac{1}{4} \left( \ln \frac{4}{k!} - 1 \right) k^{2} + O(k^{4} \ln k!),$$

$$E(k) = 1 + \frac{1}{2} \left( \ln \frac{4}{k!} - \frac{1}{2} \right) k^{2} + O(k^{4} \ln k!),$$

$$k' = \sqrt{(1-k^{2})}, k + 1, k' + 0. \tag{4.11}$$

On substituting these expansions in (4.9), we obtain for  $\lambda \epsilon_n^0 \rightarrow 0$ ,

$$\Delta E^{\text{I/KB}} \simeq \frac{2\sqrt{2}}{\pi} \left( 1 + 0 \left( \lambda \epsilon_n^0 \right) \right) \exp \left[ -\frac{\sqrt{2}}{3\lambda} \left( 1 + 0 \left( \lambda \epsilon_n^0 \ln \lambda \epsilon_n^0 \right) \right]. \tag{4.12}$$

LABIL (IV.1) ; the Gevole minimum escillator eigenvalues for various velues of  $\lambda$ . The values table is tabulated are  $\epsilon_n(\lambda) = \Xi_n(\lambda) + \frac{1}{\lambda}$ , where  $\Xi_n(\lambda)$  are the eigenvalues of  $\Xi(1,\lambda)$ .

· C.	1.582 501 444 655 76	4,005 049 195 465 72 4,006 655 456 749 52	5.347 173 384 932 61 6 376 869 564 055 15	3.162 324 720 541 12 3.514 357 705 554 85
. 62	1.595 527 585 C44 2 1.395 527 587 151 C	4.092 (25 112 82( 5 4.052 628 608 428 7	6.540 484 653 304 1 6 640 553 622 517 3	5.CC3 118 554 271 2 5.CC5 576 381 162 1
. 61	1.464 45 505 297 7	4,170 153 605 595 5	6.870 088 855 714 0 6.870 088 355 714 0	9.498 578 587 197 8 9.458 578 587 191 1
~	0 0	n n	4 rv	. 9

ر ر	уn.	Ç.).	.05
		- A representation of the confidence of the conf	
S	1.371 122 256 557 54	1.558 422 165 747 79	1.345 027 201 596 16
Н	1,371 500 461 512 93	1,350 133 557 773 29	1.550 528 987 540 87
2	5.561 359 951 815 14	5.746 917 560 727 93	5.542 342 543 859 88
10	3,518 267 537 997 13	5.848 838 500 057 5)	5.515 6C6 255 UJ8 87
4	5.838 \$11 090 504 90	5.359 059 360 284 71	5.181 424 577 100 54
ғv	6,185 906 203 845 25	6,177 383 138 505 23	235 545
9	7.424 039 289 557 84	7.470 115 938 266 45	7.750 946 881 292 35
7	8,509 274 635 240 75	8.849 281 200 220 76	751 706 581

TABER (IV.1)(... Contd.) The Couble Liminum cacillator eisenvalues for various values of A.

TABID (IV.1) (...Confd.): The double minimum oscillator eigenvalues for various values of A.

n A	, 0.7	.68	
0	1,525 374 074 208 55	1,298 249 887 534 75	1.268 237 534 205 14
<b>-</b> I	1.745 575 516 287 57	1.340 294 971 351 90	1.341 520 024 457 36
2	3.342 216 720 258 57	3.184 662 4+3 124 35	3.075 954 555 741 08
23	5.875 1.29 937 507 83	3,881 19v 140 võc 9E	5.554 COI 892 576 CA
ý.	5.127 369 933 494 84	5.288 919 C12 085 5u	5.455 992 042 717 06
2	6 472 534 E6v 4v2 36	5.787 428 109 527 17	7.059 115 211 828 7.1
9	8,137 40; 157 792 95	8,488 978 765 420 45	S.875 201 963 171 05
7	9,826 274 4.15 911 45	16.340 161 915 554 6	1C.844 973 404 181 C

TABEE (IV.1)(...Contd.); The Couble minimum cscillator eigenvalues for various values of ).

~/	,10	51.	.20
ؽ	1.23. 5C7 152 785 Cu	1.052 455 247 355 45	C 541 750 342 c76 66
<del> </del>	1,546 gáu 363 922 54	956 J	550 214 055
2	5,403 438 545 456 20	3.053 667 276 570 65	5.270 577 201 715 50
10	4.0r3 546 6.0 767 6u	4,585.838 495 343 43	5.148 274 740 0,5 02
4	5.605 133 795 683 15	6.516 658 311 767 34	7.567 134 678 671 73
Ľርን	7,556 113 319 664 21	8,564 042 111 907 84	5.827 517 529 946 06
9	9.256 331 577 195 00	11.008 558 005 345 1	12.497 859 524 785 5
7	11.354 728 355 395 9	13.526 C58 520 997 8	15.553 312 548 653 9

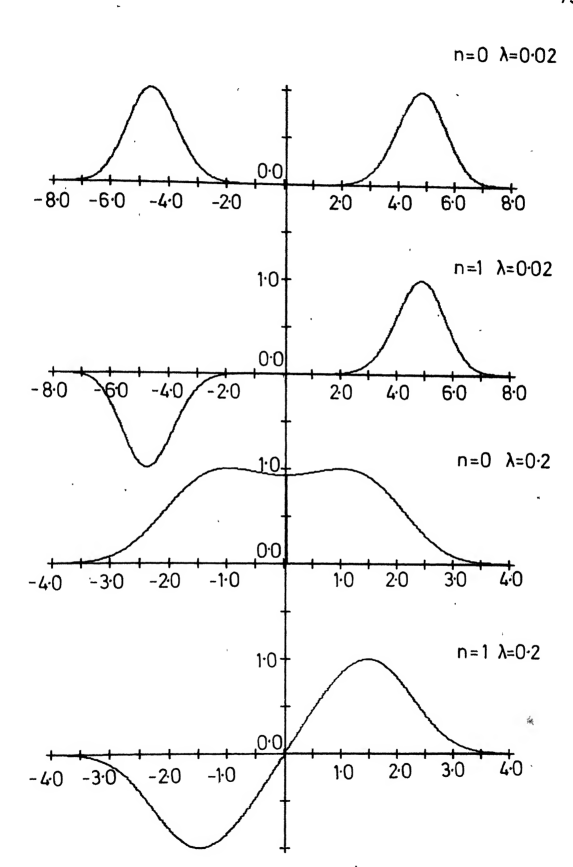


Fig. IV-1 Double Minimum Oscillator Eigenfunctions

MABLE (IV.2): Comparison of the WkB values for the splitting with the corresponding accurate values

λ	ΔE accurate†	$\Delta E_{0,1}^{accurate}/\Delta E_{0,1}^{WKB}$
.02	0.000 000 002 107 2	1.080
(13)	0.000 004 339 777 6	1.082
. 0.1	0.000 186 225 055 5	1.085
.05	0.001 711 494 025 5	1.084
• .06	0.007 299 775 950 8	1.084
.07	U.U?U U91 542 078 8	1,082
.08	0.042 045 083 827 6	1.077
.09	0.073 282 440 234 2	1.069
.10	0.112 433 706 136 5	1.048
. 1.1.	0.157 407 347 591 1	1.048
.12	0.206 073 033 949 6	1.035
.13	0.256 645 277 892 3	1.022
.14	U.307 785 906 563 0	1.012
.15	0.358 587 642 582 8	J 004
.16	0.408 462 474 699 9	1.003
. 1.7	0.457 059 973 642 5	1.010

 $<sup>^{\</sup>dagger} \Delta \mathbb{H}_{\mathcal{O}, \mathcal{I}} = \mathbb{H}_{\mathcal{I}}(\lambda) - \mathbb{H}_{\mathcal{O}}(\lambda).$ 

#### CHAPTER V

#### THE TRANSITION MOMENTS

tor energy eigenstates  $|n\rangle$  and  $|n'\rangle$  are the matrix elements  $\langle n|\chi^k|n'\rangle$ . The best known estimates (Chan and Stelman 1963, Loid 1970) of the transition moments were obtained for the pure quartic oscillator using variational eigenfunctions. The variational eigenfunctions are known to be such less accurate than the corresponding eigenvalues and are unsuitable for the computation of the transition moments of high accuracy. The occurate evaluation of the eigenvalues and the eigenvalues in this work makes it possible to obtain accurate transition moments. Further, the matrix elements  $\langle n|\chi^k|n'\rangle$  satisfy an exact linear recurrence relation in the index k (Generjee 1977). The recurrence relation makes possible the evaluation of all higher moments in terms of lover moments and eigenvalues, without integration.

# V.1 The Recurrence Relation in < n | xk | n'>

c consider the class of Hamiltonians  $H=p^2+V(x)$ , where V(x) is a real polynomial function and  $H|n>=E_n|n>$ . Then

$$\langle n | [H, W] | n' \rangle = (E_n - E_n) \langle n | W | n' \rangle$$
 (5.1)

for an arbitrary operator W. On setting  $V = x^k$  and  $x^{k-1}p$  successively in (5.1) and using the Schrödinger equation, following relations are obtained:

$$(E_{n} - E_{n}) < n | x^{k} | n^{r} > = -k(k-1) < n | x^{k-2} | n^{r} > - 2ik < n | x^{k-1} p | n^{r} >$$

$$(5.2)$$

$$(E_{n} - E_{n}) < n | x^{k-1} p | n^{r} > = -(k-1)(k-2) < n | x^{k-3} p | n^{r} >$$

$$- 2i(k-1) | E_{n} < n | x^{k-2} | n^{r} >$$

$$+ 2i(k-1) < n | x^{k-2} V(x) | n^{r} > + i < n | x^{k-1} V^{r}(x) | n^{r} > ,$$

$$\text{where } V^{r}(x) = \partial V(x) / \partial x.$$

$$(5.3)$$

Eliminating the matrix elements of  $\mathbf{x}^{k-1}\mathbf{p}$  and  $\mathbf{x}^{k-3}\mathbf{p}$  from the above two relations, we obtain

$$4k(k-1) < n | x^{k-2} V | n' > + 2k < n | x^{k-1} V' | n' >$$

$$= (\mathbb{E}_{n} - \mathbb{E}_{n'})^{2} < n | x^{k} | n' > + 2k(k-1)(\mathbb{E}_{n} + \mathbb{E}_{n'}) < n | x^{k-2} | n' >$$

$$+ k(k-1)(k-2)(k-3) < n | x^{k-4} | n' > .$$
(5.4)

For a polynomial potential V(x), the equation (5.4) reduces to the required recurrence relation. The number of initial matrix elements required to start the recursion is  $\mu$  for a polynomial potential V(x) of degree  $2\mu$ . The recursion involves the same elements of all the matrices. Thus the nn'-tk'

element of the lowest  $\mu$  matrices and the two eigenvalues  $\mathbb{F}_n$  and  $\mathbb{F}_n$ , are sufficient to determine the nn'-th element of  $\mathbb{F}_n$  for any  $\mathbb{F}_n$ . It may be noted that without the above recursion one needs all the matrix elements of the matrix  $\mathbb{F}_n$  to determine a single element of  $\mathbb{F}_n$ . In the case of the cuartic anharmonic (potential function  $\mathbb{F}_n$ ) the relation and the pure quartic oscillator  $(\mathbb{F}_n)$  the relation (1.4) yields the following recurrence relations respectively:

$$4!(k+1) \lambda < n | x^{k+2} | n' > = [(E_n - E_{n'})^2 - 4k^2] < n | x^{k} | n' > +$$

$$2k(k-1)(E_n + E_{n'}) < n | x^{k-2} | n' >$$

$$+ k(k-1)(k-2)(k-3) < n | x^{k-4} | n' > \lambda$$
(5.5)

and

4h(k+1) 
$$\lambda < n | x^{k+2} | n' > = (E_n - E_{n'})^2 < n | x^{k+2} | n' >$$
+  $2k(k-1)(E_n + E_{n'}) < n | x^{k-2} | n' >$ 
+  $k(k-1)(k-2)(k-3) < n | x^{k-4} | n' > x$  (5.6)

Thus all higher moments for any particular transition of the quartic anharmonic of the pure quartic oscillator may be obtained in terms of the lowest non-zero moment of that transition. The initial requirements in these cases is effectively reduced to one moment because of the even symmetry of the Hamiltonian, the other moment being zero. In Table (V.1) the ratio of the Ul element for the matrices x,  $x^2$ , ...  $x^{25}$  for the quartic anharmonic and the pure quartic oscillators are tabulated. For large k the recursions (5.5) and (5.6) yield

$$\lambda < n | x^{lt} | n' > / < n | x^{lt-4} | n' > \sim k^2, \quad k \to \infty , \qquad (5.7)$$

which describes the asymptotic behaviour of the moments in these cases.

# 4.2 Computation of the Transition Moments

The lower moments between various pairs of the anharmonic oscillator eigenstates may now be computed in the following manner. The expansion coefficients  $\{a_m(E)\}$  for the required eigenfunctions are evaluated by substitution the corresponding computed eigenvalues in the recursion for  $\{a_m(E)\}$ . We include as many number of coefficients  $\{a_m\}$  in the expansion of an eigenfunction as were required for obtaining the eigenvalue stable to 16 significant figures. The range of integration for the evaluation of the transition moments is truncated at  $x = x_A$ , the point in the nonclassical region at which the computed eigenfunction just begins to increase in magnitude. Since the eigenvalues used are accurate to 15 significant figures, the computed

cigenfunctions reach extremely small values in the nonclassical region before they finally start increasing in magnitude for large |x| (see section II.5). The contribution to the transition moments from the rest of the configuration space is estimated to be  $<0(10^{-14})$ . Thus,

$$< n \mid x^{k} \mid n' > = \sum_{j} \sum_{j'} a_{j}^{(n)} a_{j'}^{(n')} \int_{-x_{A}}^{+x_{f_{1}}} x^{k+j+j'} e^{-\{\alpha^{(n)} + \alpha^{(n')}\}x^{2}} dx$$

$$(5.8)$$
where  $\mid n > = \frac{1}{\sqrt{N}} \sum_{j} a_{j}^{(n)} e^{-\alpha(n)x^{2}}, \mid n' > = \frac{1}{\sqrt{N}} \sum_{j'} a_{j'}^{(n')} e^{-\alpha(n')}x^{2}$ 

If and N' are the normalization constants for the respective state the integrals involved in (5.8) may be expressed in terms of the integrals  $I_{2s}(\beta) = \int\limits_0^1 x^{2s} e^{-\beta x^2} dx$ , (s = integer), which satisfy the recurrence relation (see Appendix B),

$$(2s-1) I_{2s-2}(\beta) - (2s+1+2\beta) I_{2s}(\beta) + 2\beta I_{2s+2}(\beta) = 0.$$

$$(5.9)$$

The actual procedure for the evaluation of  $I_{2s}(\beta)$  is described in Appendix B. The computation of moments therefore requires no integration.

The non-zero matrix elements of x and  $x^2$  in the lowest ten eigenstates of the guartic anharmonic and the pure quartic oscillators for  $\lambda = 1$  were thus computed and are presented in Tables (V.2) and (V.3). Each moment given in the Tables

(V.2) and (V.3) has been checked by varying  $\alpha$  in the appropriate range and is claimed to be accurate to all 12 figures given in tables. The transition moments for the quartic anharmonic oscillator are reported for the first time in this work. Among the earlier literature only a few non-zero noments for the pure quartic oscillator were evaluated to some accuracy (Chan and Stelman 1965, feid 1970). The corresponding present values for the pure quartic oscillator moments are at least 5-6 figures more accurate. Further, the recursions (0.5) and (5.6) give all the higher moments to the same accuracy as of the lowest non-zero moment for that transition, without integration.

\* \* \* \* \*

The work presented in this thesis forms a part of a paper entitled 'The Anharmonic Oscillator' accepted for publication in the Proceedings of the Royal Society.

TABLE (V.1): The ratio  $[\langle 1|x^k|0\rangle/\langle 1|z|0\rangle]$  for the Quartic Anharmonic and the Pure Quartic Oscillators from recursions (5.5) and (5.6).

lc	[<1 xk 0><1	x[0>]
15	The Pure Quartic Oscillator $(\lambda = 1)$	The Quartic Anharmonic Uscillator $(\lambda = 1)$
•	-	
].	1.0	1.0
ÿ	0.937 978 052 782 871	0.825 567 331 595 526
5	1.301 642 584 655 12	1.073 506 435 204 95
7	2.604 681 483 536 68	1.362 740 950 928 00
C	6.036 284 029 521 49	3.986 125 267 322 52
11	16.628 197 346 910 1	10.048 610 500 904 6
15	51.631 352 902 977 5	28.965 590 251 421 1
15	178.629 006 811 098	93.500 722 956 452 3
17	678.615 442 628 609	332.840 550 182 697
10	2 799.464 881 619 44	1 291.298 361 789 83
23.	12 430.058 107 522 6	5 409.354 773 052 46
25	58 981.717 084 750 C	24 284.238 551 754 3
25	297 337.007 041 684	116 109.487 781 311

The Nonzero Matrix Elements <n|x|n'> and <n $|x^2|n'>$  between the Lowest len bigenstates of the Pure Justic Oscillator  $(E=v^2+\lambda x^{\frac{A}{2}},\lambda=1)$ . ↑ABLB ( ∵.2 ) :

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TABLE (V.3) : The Fonzero Latrix Elements  $\langle n|x|n'\rangle$  and  $\langle n|x^2|n'\rangle$  between the Lovest Ten Eigenstables of the Quartic Anharmonic Oscillator (E =  $p^2+x^2+\lambda x^4$ ,  $\lambda$  = 1).

TABLE (V.3)(...Contd.)

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4	0.036 182 636 892 -0.750 963 812	-0.750 963 812 311	157 505 501 751		
9	-0.002 348 407 864	0.065 980 790	544 -1.080 139 164 215	2,251 705 958 855	
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5		1.467 523 215 391	-0.936 356 744 621	0.078 572 602 359	-0.004 552 669 142	The second secon
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#### APPENDIX A

We describe here the procedure adopted for obtaining initial estimates of the eigenvalues in the 'boundary layer' when n is large. Let us suppose that the decired eigenvalue is  $E_n$  ( $\lambda = \lambda_b$ ), where  $(n, \lambda_b)$  lies in the 'boundary layer'. An initial estimate for it is obtained as follows:

- (i) We first obtain accurate eigenvalue  $\mathbb{E}_n$  ( $\lambda=\lambda_0$ ), where  $(n,\,\lambda_0)$  lies in the pure anharmonic region and the initial estimate for it is obtainable from the WKB formula.
- (ii) The desired value  $\lambda = \lambda_b$  is reached through a sequence of intermediate values  $\{\lambda_i\}$ ,  $i=0,1,2,\ldots$ . The initial estimate for  $E_n$  ( $\lambda = \lambda_{i+1}$ ) is obtained using accurate values of  $E_n$  ( $\lambda = \lambda_i$ ) and the Taylor series expansion. It gives

$$\mathbb{E}_{n}^{\text{initial}} \left( \lambda_{\underline{i}+\underline{1}} \right) = \mathbb{E}_{n} \left( \lambda_{\underline{i}} \right) + \left( \lambda_{\underline{i}+\underline{1}} - \lambda_{\underline{i}} \right) \left. \frac{\partial \mathbb{E}_{n}}{\partial \lambda} \right|_{\lambda = \lambda_{\underline{i}}}, \quad \underline{i}=1,2,\ldots$$

where

$$\frac{\partial E_n}{\partial \lambda} \Big|_{\lambda = \lambda_i} \simeq \frac{E_n(\lambda_i) - E_n(\lambda_{i-1})}{\lambda_i - \lambda_{i-1}}$$

The values  $\{\lambda_i\}$  are chosen sufficiently close to each other so that  $E_n(\lambda_{i+1})$  may be computed avoiding jumps to  $E_{n+1}(\lambda_{i+1}).$ 

(iii)  $E_n^{initial}(\lambda_{i+1})$  is refined to 15-figure accurate eigenvalue  $E_n(\lambda_{i+1})$  by the method used in this thesis (Section II.2). (iv) Steps (ii) and (iii) are continued till the value  $\lambda = \lambda_b$  is reached.

#### APPENDIX B

The recursive evaluation of the integrals defined by

$$I_{2s}(\beta) = \int_{0}^{1} x^{2s} e^{-\beta x^{2}} dx$$
 (B.1)

i. considered in this Appendix. On integrating (B.1) by parts, we obtain a following inhomogeneous recurrence relation

$$2\beta I_{2s+2}(\beta) - (2s+1) I_{2s}(\beta) + e^{-\beta} = C.$$
 (B.2)

Hewriting (B.2) on replacing the index 's' by 's-2'

$$2\beta I_{2s}(\beta) - (2s-1) I_{2s-2}(\beta) + e^{-\beta} = 0$$
 (B.3)

and eliminating the inhomogeneous part from (B.2) and (B.5), one obtains a 3-term homogeneous recurrence relation for  $I_{2s}(\beta) \ :$ 

$$2\beta I_{2s+2}(\beta) - (2s+1+2\beta) I_{2s}(\beta) + (2s-1) I_{2s-2}(\beta) = 0.$$
(B.4)

The integrals  $I_{2s}(\beta)$  may therefore be computed for any value of s by successive application of the relation (B.4). However, since the computations are carried out perforce with rounded values, the relative errors grow and overtake the wanted function when a straightforward use of the above recursion

is made in forward direction (increasing s). This occurs when a lecurrence relation has two independent solutions and the solution desired is disnishing as the index 's' increases, while the compunion solution is increasing. reversing the direction the roles of the two solutions are interchanged and the contribution of desired solution now increases while the un anted solution dimniphes (Abramowitz and Stegun 1965). Computation of the integrals  $I_{2s}(\beta)$  is therefore done by applying the recursion (1.4) in backward direction (decreasing s). The recursion is started from a cufficiently higher index 's' using arbitrary starting ( Miller 1952). The values obtained in this values manner differ from the desired solution by a constant multiplier which is calculated from the values of  $I_{\Omega}(\beta)$ obtained from the tables for the error functions.

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